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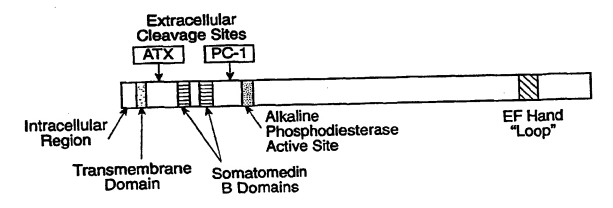
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#### (57) Abstract

The present invention relates, in general, to autotaxin. In particular, the present invention relates to a DNA segment encoding autotaxin; recombinant DNA molecules containing the DNA segment; cells containing the recombinant DNA molecule; a method of producing autotaxin; antibodies to autotaxin; and identification of functional domains in autotaxin.

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AUTOTAXIN: MOTILITY STIMULATING PROTEIN USEFUL IN CANCER DIAGNOSIS AND THERAPY

This application is a continuation-in-part of application serial no. 08/249,182 filed May 25, 1994, which is a continuation-in-part of application serial no. 07/822,043 filed on Jan. 17, 1992.

# Field of the Invention

The present invention relates, in general, to a motility stimulating and compositions comprising the same. In particular, the present invention relates to a purified form of the protein and peptides thereof, for example, autotaxin (herein alternative referred to as "ATX"); a DNA segment encoding autotaxin; recombinant DNA molecules containing the DNA segment; cells containing the recombinant DNA molecule; a method of producing autotaxin; antibodies to autotaxin; and methods of cancer diagnosis and therapy using the above referenced protein or peptides thereof and DNA segments.

# Background of the Invention

Cell motility plays an important role in embryonic events, adult tissue remodeling, wound healing, angiogenesis, immune defense, and metastasis of tumor cells (Singer, 1986). In normal physiologic processes, motility is tightly regulated. On the other hand, tumor cell motility may be aberrantly regulated or autoregulated. Tumor cells can respond in a motile 25 fashion to a variety of agents. These include hostderived factors such as scatter factor (Rosen, et al., 1989) and growth factors (Kahan, et al., 1987; Stracke, et al.; Tamm, et al., 1989; Wang, et al. 1990; and Jouanneau, et al. 1991), components of the extracellular matrix (McCarthy, et al. 1984), and tumor-secreted or autocrine 30 factors (Liotta, et al. 1988; Ruff, et al. 1985; Atnip, et al. 1987; Ohnishi, et al. 1990; Silletti, et al. 1991; and Watanabe, et al. 1991).

Many types of host-derived soluble factors act

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in a paracrine fashion to stimulate cell locomotion. Motility-stimulating proteins called "scatter factors" have been identified which are produced by embryonic fibroblasts and by smooth muscle cells (Stoker, et al. 1987). Scatter factors stimulate random and directed 5 motility by epithelial cells, keratinocytes, vascular endothelial cells and carcinoma cells (Stoker, et al. 1987; Rosen, et al. 1990; and Weidner, et al. 1990), but not fibroblasts. In addition, a number of host-secreted growth factors have been demonstrated to stimulate 10 motility in tumor cells, including nerve growth factor (Kahan, et al. 1987) insulin-like growth factor-I (Stracke, et al. 1988), interleukin-6 (Tamm, et al. 1989), interleukin-8 (Wang, et al. 1990), and acidic fibroblast growth factor (Jouanneau, et al. 1991). These paracrine 15 factors may influence "homing" or the directionality of tumor cell motility.

In contrast to these host-derived factors, many types of tumor cells have been found to produce proteins termed "autocrine motility factors" which stimulate motility by the same tumor cells which make the factor (Liotta, et al. 1986). Autocrine motility factors are not specific for a given type of cancer cell but have a wide spectrum of activity on many types of cancer cells (Kohn, et al. 1990), with little effect on normal fibroblasts or leukocytes.

Autocrine motility factors identified to date act through cell-surface receptors (Stracke, et al. 1987; Nabi, et al. 1990; Watanabe, et al. 1991) resulting in pseudopodial protrusion (Guirguis, et al. 1987) leading to both random and directed migration (Liotta, et al. 1986; Atnip, et al. 1987; Ohnishi, et al. 1990).

Prior studies of human A2058 melanoma cells have demonstrated that these cells are a particularly rich source of autocrine motility factors. An autocrine motility factor with a molecular mass of approximately 60

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kDa has been previously isolated from the conditioned media of these cells. (Liotta, et al. 1986). Similar tumor cells derived or induced factors with the same molecular weight have subsequently been reported and purified by several investigators (Atnip, et al. 1987; Schnor, et al. 1988; Ohnishi, et al. 1990; Silletti, et al. 1991; Watanabe et al. 1990). Such factors are thought to play a key role in tumor cell invasion.

Most of the motility factors identified to date have not been purified to homogeneity and have not been sequenced. The novel tumor motility factor of the present invention, named herein as autotaxin ("ATX"), has been purified and verified to be a homogeneous sample by twodimensional gel electrophoresis. The protein of the present invention is unique from any previously identified or purified motility factor. The molecular size of ATX is about 125 kilo Daltons ("kDa") and it has an isoelectric point of approximately 7.7. ATX stimulates both random and directed migration of human A2058 melanoma cells at picomolar concentrations. The activity of the ATX factor is completely sensitive to inhibition by pertussis toxin. No significant homology has been found to exist between the protein of the invention and any mammalian protein including previous factors known to stimulate cell motility.

There is a great clinical need to predict the aggressiveness of a patient's individual tumor, to predict the local recurrence of treated tumors and to identify patients at high risk for development of invasive tumors. The present invention provides a functional marker which is functionally related to the invasive potential of human cancer. The invention further provides an assay for this secreted marker in body fluids, or in tissues. The assay of the invention can be used in the detection, diagnosis, and treatment of human malignancies and other inflammatory, fibrotic, infectious or healing disorders.

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## SUMMARY OF THE INVENTION

The present invention relates, generally, to a motility stimulating protein and corresponding peptides thereof, and to a DNA segment encoding same. A human cDNA clone encoding a tumor cell motility-stimulating protein, herein referred to as autotaxin or "ATX", reveals that this protein is an ecto/exoenzyme with significant homology to the plasma cell membrane differentiation antigen PC-1. ATX is a 125 kDa glycoprotein, previously isolated from a human melanoma cell line (A2058), which elicits chemotactic and chemokinetic responses at picomolar to nanomolar concentrations.

It is a specific object of the present invention to provide autotaxin and peptide fragments thereof.

It is a further object of the present invention to provide a DNA segment that encodes autotaxin and a recombinant DNA molecule comprising same. It is a further object of the present invention to provide a cell that contains such a recombinant molecule and a method of producing autotaxin using that cell.

Another object of the present invention is the identification of a transmembrane domain of the human liver autotaxin protein and its apparent absence in tumorous forms of autotaxin. The tumorous form of autotaxin appears to be a secreted protein. The present invention relates to utilization of the different sites of localization for diagnosis and prognosis of the stages of tumor progression. Further, the invention relates to treatment methods, designed to advantageously block the secreted form of autotaxin activity while having little effect on the membrane-bound form of autotaxin.

Yet another object of the present invention relates to the identification of a highly variable region within the autotaxin gene, called a "hot spot". The variations in sequence apparently result in mutations, insertions, deletions and premature termination of

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translation. The present invention relates to manipulating this region so as to alter the activity of the protein. Further, the hot spot can serve as a marker in tumor diagnosis differentiating between different forms of the autotaxin protein.

It is yet another object of the present invention to provide a method of purifying autotaxin.

It is a further object of the present invention to provide cloned DNA segments encoding autotaxin and fragments thereof. The cDNA encoding the entire autotaxin protein contains 3251 base pairs, and has an mRNA size of approximately 3.3 kb. The full-length deduced amino acid sequence of autotaxin comprises a protein of 915 amino acids. Database analysis of the ATX sequence revealed a 45% amino acid identity (including 30 out of 33 cysteines) with PC-1, a pyrophosphatase/type I phosphodiesterase expressed on the surface of activated B cells and plasma cells. ATX, like PC-1, was found to hydrolyze the type I phosphodiesterase substrate p-nitrophenyl thymidine-5'monophosphate. Autotaxin now defines a novel motility-regulating function for this class of ecto/exo-enzymes.

Further objects and advantages of the present invention will be clear from the description that follows.

# BRIEF DESCRIPTION OF THE DRAWINGS

25 Figure 1. Fractionation of ATX by hydrophobic interaction. A 200 ml sample of A2058 conditioned media was chromatographed on a 200 mL column of phenyl Sepharose-4B. Buffer A was 50 mM Tris (pH 7.5), 5% methanol, and 1.2 M ammonium sulfate. Buffer B was 50 mM Tris (pH 7.5), 5% methanol and 50% ethylene glycol. The gradient (----) represents a double linear gradient with decreasing ammonium sulfate (1.2 to 0.0 M) and increasing ethylene glycol (0 to 50%). Absorbance was monitored at 280 nm (----) and indicated that most of the protein did not bind to the column. Ten ml fractions were assayed for

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motility stimulating capacity using the Boyden Chamber assay (o). The peak of motility activity occurred between 900 and 1050 minutes, ~ 12% of the gradient.

Figure 2. Isolation of ATX by lectin affinity chromatography. 20 ml portions of the phenyl Sepharose activity peak were affinity purified on a 40 ml Concanavalin A Affi-Gel column. The bound components were eluted with a step gradient (----) of methyl  $\alpha$ -Dmannopyranoside (0.0 mM, 10 mM, and 500 mM) in a buffer consisting of 0.05 M Tris (pH 7.5), 0.1 M NaCl, 0.01 M CaCl<sub>2</sub> and 20% ethylene glycol. Absorbance was monitored at 280 nm (\_\_\_\_\_) and indicated that the majority of the protein components did not bind to the column. Motility was assayed in 10 mL fractions (...o...) and was found predominantly in the 500 mM elution concentration. 15 One of seven chromatographic runs is shown.

Purification of ATX by weak anionic Figure 3. exchange chromatography. Approximately 30% of the activity peak eluted from the Con A affinity column was applied to a ZORBAX BioSeries-WAX column. The bound components were eluted with an NaCl gradient (----) in a buffer consisting of 10 mM Tris (pH 7.5) and 30% ethylene glycol. Motility (o) was assayed in 1.0 ml fractions. The peak of activity eluted in a discrete but broad region in the shallow portion of the gradient. Absorbance was monitored at 230 nm (\_\_\_\_\_). The majority of the protein components not associated with activity bound strongly to the column were eluted at 1.0 M NaCl. One of two chromatographic runs is shown.

Figure 4. Purification of ATX by molecular sieve exclusion chromatography. The entire activity peak eluted from the weak anion exchange column was applied to a series of TSK columns (4000SW, 4000SW, 3000SW, and 2000SW, in this order). Proteins were eluted in a buffer consisting of 0.1M NaPO4 (pH 7.2) with 10% methanol and 10% ethylene glycol. Two major protein peaks were evident

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by monitoring the absorbance at 235 nm (\_\_\_\_\_). Motility (...o...) was assayed in 0.4 ml samples and found predominantly in the first, smaller, protein peak.

anionic exchange chromatography. Approximately 15% of the activity peak from the molecular sieve exclusion series was applied to a Pro-Pac PA1 column. Protein which bound to the column was eluted with a NaCl gradient (----) in a buffer consisting of 10 mM Tris (pH 7.5), 5% methanol and 20% ethylene glycol. Absorbance was monitored at 215 nM (\_\_\_\_). Motility activity was assayed in 1.0 ml fractions at two different dilutions: 1/5 (...o...) or 1/15 (.\_\_.o.\_\_.). Activity was found to correspond to a double protein peak in the central region of the gradient.

Figure 6A, 6B and 6C. Protein components associated with the activity peaks from various stages of purification. The activity peak from each chromatographic fractionation was pooled, concentrated and analyzed by SDS-polyacrylamide gel electrophoresis. Molecular weight standards are in Lane 1 for each panel. Panel 6A) 8-16% gradient gel of the first three purification steps, run under non-reducing conditions. Lane 2 is an aliquot of the pooled activity peak eluted from the phenyl sepharose fractionation. Lane 3 is an aliquot of the pooled activity peak eluted from the Con A affinity purification. Lanes 4 and 5 show the "peak" and "shoulder" of activity fractionated by weak anion exchange chromatography (Figure Panel 6B) 7% gel of the activity peak fractionated by molecular sieve exclusion chromatography. Lanes 2 and 3 show the protein separation pattern of the total pooled activity peak when the gel was run under non-reducing and reducing conditions, respectively. Panel 6C) 8-16% gradient gel of the final strong anionic exchange chromatographic separation, run under non-reducing conditions. Lane 2 comprises ~1% of the total pooled activity peak eluted from the column.

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Figure 7. Two-dimensional gel electrophoresis of ATX. Purified ATX (Figure 6, Panel C) was subjected to non-equilibrium isoelectric focusing (5 hr. at 500v), then applied to a 7.5% SDS-polyacrylamide gel for the second dimension. The pH separation which resulted was measured in 0.5 cm samples of concurrently run tube gels and is shown at the top. Molecular weight standards for the second dimension are shown on the right. This analysis reveals a single component with pI =  $7.7 \pm 0.2$  and M = 120,000.

10 Figure 8. Dilution curve of ATX. Purified ATX (Figure 6, Panel C) was serially diluted and tested for motility-stimulating activity. The result, with unstimulated background motility subtracted out, shows that activity is half-maximal at ~ 500 pM ATX.

Figure 9. Pertussis toxin (PT) sensitivity of ATX. A2058 cells were pre-treated for 1 hr. prior to the start of the motility assay with 0.5  $\mu$ g/ml PT in 0.1% BSA-DMEM or with 0.1% BSA-DMEM alone (for untreated control). The motility activity stimulated by purified ATX (Figure 6, Panel C) was then assessed for the two treatment groups. The result, expressed as cells/HPF  $\pm$  S.E.M. with unstimulated background motility subtracted out, reveals profound inhibition of PT-treated cells (hatched) compared to untreated cells (solid). PT had no effect on cell viability. S.E.M.'s were < 10%.

Figure 10. Checkerboard analysis of ATX-stimulated motility. Varying dilutions of autotaxin were added to the upper chamber with the cells and/or to the lower chamber, as shown. Motility response, expressed as cells/HPF  $\pm$  S.E.M., was assessed for each point in the checkerboard.

Figure 11. Purification of ATX peptides on HPLC. ATX, purified to homogeneity by strong anionic exchange chromatography, was sequentially digested by cyanogen bromide, subjected to reduction and

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pyridylethylation, and digested by trypsin. The resulting peptides were purified on an Aquapore RP300 C-8 reverse phase column using a (0-70)% acetonitrile gradient in 0.1% trifluoroacetic acid (----). The absorbance was monitored at 215 nm (\_\_\_\_\_) and peaks were collected. Seven peaks, chosen at random for N-terminal amino acid sequence analysis, are shown with appropriate numbers.

Figure 12. Cloning Strategy, schematically depicted.

Figure 13. Schematic Diagram of autotaxin gene region.

For A2058: 4C11 is the original DNA clone obtained by screening an A2058 cDNA expression library in λgt11 with anti-peptide ATX-102. Upstream ATX peptide sequences were utilized for PCR amplification of A2058 mRNA, using the technique of reverse transcription/PRC. These peptides include ATX-101, ATX-103, and ATX-224. The approximate localization of each of peptide was obtained by matching the peptide with its homologous region on PC-

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For N-tera 2D1, a λgt1 0 cDNA library was amplified and the cDNA inserts were isolated. PCR amplification, based on homologies with A2058 sequence, was utilized for DNA sequencing.

For normal human liver, a mRNA from liver was amplified with 5'RACE using primers from the known ATX-224 region of A2058 and N-tera 2D1.

Figure 14. Schematic match-up of ATX peptides with putative A2058 protein sequence.

Figure 15. Schematic match-up of ATX peptides with putative N-tera 2D1 protein sequence.

Figure 16: ATX Treatment with PGNase F.

Partially purified ATX was treated with 60 mU/ml PNGase F
at 37°C for 16 hr under increasingly denaturing
conditions. The treated ATX samples were separated by SDS
polyacrylamide gel electrophoresis run under reducing

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conditions and stained with Coomassie blue G-250. Lane 1 contains untreated ATX (arrow) with no enzyme added. Lane 2 contains the reaction mixture run under non-denaturing conditions (50 mM tris/10% ethylene glycol, pH 7). Lanes 3 and 4 have added 0.1 M  $\beta$ -mercaptoethanol and 0.5%

Nonidet-P40, respectively. Lanes 5 and 6 contain the reaction mixtures in which the ATX sample was first boiled for 3 min in 0.1% SDS with (lane 6) or without (lane 5) 0.1 M  $\beta$ -mercaptoethanol, then had 0.5% Nonidet-P40 added to prevent enzyme denaturation. The enzyme can be detected as an ~44 kDa band in lanes 2-6.

Figure 17: Effect of varying concentrations of PNGase F on ATX molecular weight and motility-stimulating activity. Partially purified ATX was treated with various concentrations (range 0 - 60 mU/ml, shown on horizontal axis) of PNGase F at 37°C for 16 hr. Figure 17A shows the effect of the different treatments on ATX molecular weight. At concentrations of enzyme  $\geq$  30 mU/ml, the deglycosylation reaction appears to be complete. Figure 17B shows the effect of the identical reaction mixtures on motility-stimulating capacity (immediately below the corresponding protein band of Figure 17A). There is no significant difference between any of the treatment groups.

Figure 18: Comparison of amino acid sequences of ATX and PC-1. The amino acid sequences of ATX and PC-1 are compared. Amino acid identity is indicated by a vertical line between the sequences. The location of the putative transmembrane/signal sequence is shown by a solid line. The two somatomedin B domains are identified by dashed lines. The putative phosphodiesterase active site is indicated by emboldened lines. The loop region of a single EF hand loop region is identified with double lines. The presumed cleavage site for each protein is indicated with arrows.

Figure 19: Domain structure of ATX and PC-1.

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Putative domains are indicated for the two homologous proteins, ATX and PC-1.

# DETAILED DESCRIPTION OF THE INVENTION

Tumor cell motility is a critical component of invasion and metatasis, but the regulation of this motility is still poorly understood. At least some tumor cells secrete autocrine motility factors (AMF's) that stimulate motility in the producing cells. Like the analogous autocrine growth factors, these AMF's allow tumor cells independence from the host in this important component of the metastatic cascade. One AMF, autotaxin (ATX), has recently been purified to homogeneity from the human melanoma cell line, A2058 (Stracke, et al., 1992). The purified protein was enzymatically digested and the peptide fragments were separated by reverse phase HPLC. A number of these peptides have been sequenced by standard 15 · Edman degradation (Table 6) from different purifications and different enzymatic digestion. Sequence information, obtained initially on 19 purified tryptic peptides, confirmed that the protein is unique with no significant homology to growth factors or previously described motility factors. These peptide sequences have now been used as the basis for identifying and sequencing the cDNA clone for ATX. The present invention comprises an amino acid sequence of ATX as well as a nucleic acid sequence coding for the ATX protein.

TABLE 6. PEPTIDE SEQUENCES FOR AUTOTAXIN.

	PEPTIDE NO.	AMINO ACID SEQUENCE	SEQ ID: NO:
30	ATX-18	WHVAR	SEQ ID NO:1
	ATX-19	PLDVYK	SEQ ID NO:2
	ATX-20	YPAFK	SEQ ID NO:3
	ATX-29	PEEVTRPNYL	SEQ ID NO:5
35	ATX-34B	RVWNYFQR	SEQ ID NO:38

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•	ATX-41	HLLYGRPAVLY	SEQ ID NO:29		
	ATX-48	VPPFENIELY	SEQ ID NO:7		
	ATX-59	TFPNLYTFATGLY	SEQ ID NO:32		
	ATX-100	GGQPLWITATK	SEQ ID NO:8		
5	ATX-101/223A	VNSMQTVFVGYGPTFK	SEQ ID NO:9		
3	ATX-102	DIEHLTSLDFFR	SEQ ID NO:10		
	ATX-103	TEFLSNYLTNVDDITLVPETLGR	SEQ ID NO:11		
	ATX-104	VNVISGPIDDYDYDGLHDTEDK	SEQ ID NO:33		
	ATX-204	MHTARVRD	SEQ ID NO:39		
10	ATX-205	FSNNAKYD	SEQ ID NO:40		
	ATX-209	VMPNIEK	SEQ ID NO:41		
	ATX-210	TARGWECT	SEQ ID NO:42		
	ATX-212	(N)DSPWT(N)ISGS	SEQ ID NO:43		
	ATX-214	LRSCGTHSPYM	SEQ ID NO:44		
15	ATX-215/34A	TYLHTYES	SEQ ID NO:45		
	ATX-213/217A	AIIANLTCKKPDQ	SEQ ID NO:46		
	ATX-216	IVGQLMDG	SEQ ID NO:47		
	ATX-218/44	TSRSYPEIL	SEQ ID NO:48		
20	ATX-223B/24	QAEVSSVPD	SEQ ID NO:49		
20	ATX-224	RCFELQEAGPPDDC	SEQ ID NO:50		
	ATX-229	SYTSCCHDFDEL	SEQ ID NO:51		
	ATX-244/53	QMSYGFLFPPYLSSSP	SEQ ID NO:52		
	ATX is a glyc	osylated protein due to it	s high		
25	affinity for conca	navalin A and amino acid s	sequence		
	analysis of the ATX peptides. ATX has been demonstrated				
		coprotein whose molecular	-		
		r deglycosylation with N-g	=		
		ecular weight of the clone			
30	100kDa (secreted f	orm) or 105kDa (full lengt	h protein).		

to be a 125kDa glycoprotein whose molecular weight reduced to 100-105kDa after deglycosylation with N-glycosidase F. The calculated molecular weight of the cloned protein is 100kDa (secreted form) or 105kDa (full length protein). Based on amino acid composition, the estimated pI is 9.0 which is higher than the pI determined by 2-D gel electrophoresis analysis (7.7-8.0) of purified ATX. This difference can be explained by the presence of sialic acid residues on the sugar moieties.

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Autotaxin is secreted by A2058 human melanoma cells cultured in low abundance in serum-free conditioned Autotaxin is a potent new cytokine with molecular mass 125 kDa which has been purified to homogeneity from the conditioned medium of the human melanoma cell line, A2058, utilizing sequential chromatographic methods as described herein. This new cytokine, termed autotaxin (ATX), is a basic glycoprotein with pI ~ 7.8. ATX is active in the high picomolar to low nanomolar range, stimulating both chemotactic and chemokinetic responses in the ATX-producing A2058 cells as well as other tumor cells. This motile response is abolished by pretreatment of the cells with pertussis toxin. ATX may therefore act through a G protein-linked cell surface receptor. characteristics distinguish ATX from several small growth factors and interleukins which are implicated in tumor cell motility (Stracke et al., 1988; Ruff et al., 1985; Maciag et al., 1984; Gospodarowicz, 1984; Van Snick, 1990; Yoshimura 1987).

The protein of the present invention, which in one embodiment is derived from A2058 human melanoma cells, can be prepared substantially free from proteins with which it is normally associated using, for example, the purification protocol disclosed herein. Alternatively, the protein of the present invention can be prepared substantially free from proteins, by cloning and expressing the cDNA encoding autotaxin as disclosed herein.

A large volume of serum-free conditioned medium from appropriate producer cells (e.g., tumor cells) is collected and concentrated approximately 500-fold. This concentrated conditioned medium is then separated from other contaminating proteins by techniques that rely on the chemical and physical characteristics of the protein. These include the molecular weight, relative hydrophobicity, net charge, isoelectric focusing point,

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and the presence of lectin-binding sugar residues on the protein.

Alternatively, the protein, or functional portion thereof, can be synthesized using chemical or recombinant means.

The protein of the present invention has a potent biological activity. Purified ATX is active in the picomolar range and 1 unit of activity corresponds to a concentration of approximately 500 pM as assessed by the cell motility assay described herein and elsewhere (Stracke et al., 1989).

The protein of the present invention has a molecular size, as determined by two dimensional gel electrophoresis, of from 120 to 130 kDa, or more specifically, about 125 kDa. Further, the protein of the present invention can have a pI in the range of 7.5 to 8.0, preferably, approximately 7.7. The present invention relates to autotaxin and peptides thereof having cell motility properties as described herein. These proteins and peptides thereof can be produced by isolation from a natural host or isolation as an expression product from a recombinant host.

The present invention also relates to a DNA segment coding for a polypeptide comprising an amino acid sequence corresponding to ATX, or a unique portion of such a sequence (unique portion being defined herein as at least 5, 10, 25, or 50 amino acids). In one embodiment, the DNA segment encodes any one of the amino acid sequences shown in SEQ ID NO:1 to SEQ ID NO:11 and SEQ ID NO:26 to SEQ ID NO:33. Another embodiment uses larger DNA fragments encoding amino acid sequences shown in SEQ ID NO:34, SEQ ID NO: 36 and SEQ ID NO:38. The entire coding region for autotaxin can also be used in the present invention shown in SEQ ID NO:66 through SEQ ID NO:69.

In another embodiment, the present invention relates to a recombinant DNA molecule comprising a vector (for

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example plasmid or viral vector) and a DNA segment coding for a polypeptide corresponding to ATX, as can be prepared by one skilled in the art. Preferably, the coding segment is present in the vector operably linked to a promoter.

The present invention also relates to a recombinant protein produced from a host cell expressing a cDNA containing a coding region of ATX. Examples of ATX cDNAs from a variety of sources have been cloned and can be used for expression, including inter alia A2058 carcinoma

cells, N-tera 2D1 cells and human liver.

In a further embodiment, the present invention relates to a cell containing the above-described recombinant DNA molecule. Suitable host cells include procaryotic cells (such as bacteria, including <u>E. coli</u>) and both lower eucaryotic cells (for example, yeast) and higher eucaryotic cells (for example, mammalian cells). Introduction of the recombinant molecule into the host cells can be effected using methods known in the art.

In another embodiment, the present invention relates to a method of producing a polypeptide having an amino acid sequence corresponding to ATX. The method comprises culturing the above-described cells under conditions such that the DNA segment is expressed, and isolating ATX thereby produced.

In a further embodiment, the present invention relates to an antibody having affinity for ATX or peptide fragments thereof. The invention also relates to binding fragments of such antibodies. In one preferred embodiment, the antibodies are specific for ATX peptides having an amino acid sequence set forth in one of SEQ ID NO:1 through SEQ ID NO:11 and SEQ ID NO:26 through SEQ ID NO:34, SEQ ID NO: 36 and SEQ ID NO:38 through SEQ ID NO:52. In addition, the antibodies may recognize an entire autotaxin protein.

Antibodies can be raised to autotaxin or its fragment peptides, either naturally-occurring or recombinantly

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produced, using methods known in the art.

The ATX protein and peptide fragments thereof described above can be joined to other materials, particularly polypeptides, as fused or covalently joined polypeptides to be used as carrier proteins. In particular, ATX fragments can be fused or covalently linked to a variety of carrier proteins, such as keyhole limpet hemocyanin, bovine serum albumin, tetanus toxoid, etc. See for example, Harper and Row, (1969); Landsteiner, (1962); and Williams et al., (1967), for descriptions of methods of preparing polyclonal antisera. A typical method involves hyperimmunization of an animal with an antigen. The blood of the animal is then collected shortly after the repeated immunizations and the gamma globulin is isolated.

In some instances, it is desirable to prepare monoclonal antibodies from various mammalian hosts. Description of techniques for preparing such monoclonal antibodies may be found in Stites et al., and references cited therein, and in particular in Kohler and Milstein (1975), which discusses one method of generating monoclonal antibodies.

In another embodiment, the present invention relates to an oligonucleotide probe synthesized according to the sense or antisense degenerative sequence set forth in one of SEQ ID NO:1 through SEQ ID NO:11, SEQ ID NO:26 through SEQ ID NO:33, SEQ ID NO:39 through SEQ ID NO:52, and SEQ ID NO:55 through SEQ ID NO:65.

Protein database searches of this sequence revealed a 45% amino acid identity with the plasma cell membrane marker protein, PC-1. ATX and PC-1 appear to share a number of domains, including two somatomedin B domains, the loop region of an EF hand, and the enzymatic site of type I phosphodiesterase/ nucleotide pyrophosphatase.

Like PC-1, ATX hydrolyzes p-nitrophenyl thymidine-5' - monophosphate, a type 1 phosphodiesterase substrate. This

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enzymatic function of ATX suggests a newly identified function for ecto/exo-enzymes in cellular motility.

In a further embodiment, the present invention relates to a method of diagnosing cancer metastasis and to a kit suitable for use in such a method. Preferably, antibodies to ATX can be used in, but not limited to, ELISA, RIA or immunoblots configurations to detect the presence of ATX in body fluids of patients (e.g. serum, urine, pleural effusions, etc.). These antibodies can also be used in immunostains of patient samples to detect the presence of ATX.

In yet another embodiment, the present invention relates to in vivo and in vitro diagnostics. ATX may be radiolabelled, by means known to one skilled in the art, and injected in cancer patients with appropriate ancillary substances also known to one skilled in the art, in order to ultimately detect distant metastatic sites by appropriate imagery. The level of ATX in tissue or body fluids can be used to predict disease outcomes and/or choice of therapy which may also include ATX inhibitors.

In a further embodiment, the present invention relates to a treatment of cancer. ATX antibodies can be cross-linked to toxins (e.g., Ricin A), by means known to one skilled in the art, wherein the cross-linked complex is administered to cancer patients with appropriate ancillary agents by means known to one skilled in the art, so that when the antibody complex binds to the cancer cell, the cell is killed by the cross-linked toxin.

In another embodiment, the different localizations of the normal versus tumorous forms of the ATX proteins within the tissue can be used as a tool for diagnosis and prognosis. The stage of disease progression can be monitored by elevated levels of ATX in the extracellular space as opposed to its normal cell membranes association. In addition, treatment methods for control of tumor progression can be designed to specifically block the

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activity of the secreted form of ATX. Such methods would have a preferential effect upon secreted ATX during tumor progression while not effecting normal ATX formation.

Yet another embodiment utilizes the hot spot located in the region from approximately nucleotides 1670 through 1815, as a marker gene for identification of tissues carrying a tumorous form of ATX.

The present invention is described in further detail in the following non-limiting examples.

#### **EXAMPLES**

10 The following protocols and experimental details are referenced in the Examples that follow: Materials. The polycarbonate Nuclepore membranes and the 48-well microchemotaxis chambers were obtained from Neuro Probe, Inc. Pertussis toxin (PT), ethylene glycol 15 (biotechnology grade), methyl  $\alpha$ -D-mannopyranoside were obtained from commercial vendors. The ampholyte, pH 3-10 Bio-Lyte and pH 8-10 Bio-Lyte, were obtained from Bio-Rad. Phenyl Sepharose CL-4B; affi-Gel concanavalin A; ZORBAX BioSeries-WAX (weak anion exchange) column (9.4mm x 24cm); 20 Spherogel-TSK 4000SW, 3000SW and 2000SW columns (each 7.5mm x 30cm); the Pro-Pac PA1 (4 x 50mm) strong anion exchange column; the Aquapore RP300 C-8 reverse phase column (220 x 2.1mm); and the AminoQuant C-18 reverse phase column (200 x 2.1mm) were also obtained from 25 commercial sources.

Affi-Gel 10 affinity resin was from Bio-Rad. The GeneAmp PCR Reagent kit with AmpliTaq and the GeneAmp RNA PCR kit were purchased from Perkin-Elmer. The 5' RACE kit came from Gibco BRL Life Technologies, Inc. The p-nitrophenyl thymidine-5'monophosphate was obtained from Calbiochem Biochemicals.

Ethylene glycol (biotechnology grade) was from Fisher Biochemicals (Pittsburg, PA). Peptide N-glycosidase F ("PNGase F"), O-glycosidase, neuraminidase (Arthrobacter ureafaciens), and swainsonine ("Swn") came from

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Boehringer-Mannheim (Indianapolis, IN). Deoxymannojirimycin ("dMAN"), and N-methyl-1deoxynojirimycin ("NMdNM") were from Oxford GlycoSystems, Inc. (Rosedale, NY). Biotinylated concanavalin A, HRPconjugated streptavidin, and HRP-conjugated goat antirabbit immunoglobulin were purchased from Pierce Chemicals 5 (Rockford, IL). Polyvinyl pyrrolidone-free polycarbonate membranes and the microchemotaxis chamber were from NeuroProbe, Inc. (Cabin John, MD). Cell Culture. The human melanoma cell line A2058, originally isolated by Todaro (Todaro et al., 1980), was 10 maintained as previously described by Liotta (Liotta et al., 1986). The N-tera 2 (D1 clone) was a kind gift from Dr. Maxine Singer, Laboratory of Biochemistry, National Cancer Institute, National Institutes of Health and was maintained as described (Andrews, P.W., Goodfellow, P.N. 15 and Bronson, D.L. (1983) Cell surface characteristics and other markers of differentiation of human teratocarcinoma cells in culture.). Production of Autotaxin. A2058 cells were grown up in T-150 flasks, trypsinized, and seeded into 24,000 cm2 cell 20 factories at a cell density of 1x1010 cells/factory. After 5-6 days, the serum-containing medium was removed and the cells were washed with DPBS. The factories were maintained in DMEM without phenol red, supplemented with 4 mM glutamine, 100 units/ml penicillin, 100  $\mu$ g/ml 25 streptomycin, 5  $\mu$ g/ml crystallized bovine serum albumin, 10  $\mu$ g/ml bovine insulin, and 1  $\mu$ M aprotinin. Culture supernatants were harvested every 3 days, frozen at -40°C and replaced with fresh serum-free medium. Each cycle of supernatant was tested for ATX production with a cell 30 motility assay detailed below. Typically, a cell factory continued to be productive for 9-11 of these cycles. After accumulation of approximately 45-60 L of supernatant, the culture supernatants were thawed and concentrated down to 2-2.5 L using an Amicon S10Y30 spiral

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membrane ultrafiltration cartridge. This supernatant was further concentrated in an Amicon high performance ultrafiltration cell using Diaflo membranes. The final volume achieved from 100-200 L of conditioned medium was typically 250-400 ml. All ultrafiltrations were performed at 4°C.

Cell Motility Assays. Purification of autotaxin was monitored by testing the motility-stimulating capacity of the fractions collected from the columns. These fractions were in buffers unsuitable for a chemotaxis assay so each fraction had to be washed into an appropriate buffer, i.e., 0.1% (w/v) BSA in DPBS containing calcium and magnesium. This dialysis was performed by adding aliquots of each fraction to be tested into Centricon-30<sup>m</sup> ultrafiltration tubes, which retain molecular species larger than 30,000 daltons.

The assay to determine motility was performed in triplicate using a 48-well microchemotaxis chamber as described elsewhere in detail (Stracke et al., 1987; Stracke, et al., 1989). The Nuclepore™ membranes used in these modified Boyden chambers were fixed and stained with Diff-Quik.™ Chemotaxis was quantitated either by reading the stained membranes with a 2202 Ultroscan laser densitometer or by counting 5 randomly chosen high power fields (HPF) under light microscopy (400 x) for each replicate. Densitometer units (wavelength - 633 nm) have been shown to be linearly related to the number of cells per HPF (Taraboletti, 1987; Stracke, et al., 1989). Typically, unstimulated motility (background) corresponded to 5-10 cells/HPF and highly responding cells to 70-100 cells/HPF above unstimulated background (i.e., 75-110 total cells/HPF).

For experiments using PT, the toxin was pre-incubated with the cells for 1-2 hr. at room temperature prior to the assay and maintained with the cells throughout the assay (Stracke, et al., 1987). The treated cells were

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tested for their motility response to the chemoattractant as well as for unstimulated random motility.

Purification of Autotaxin. Ammonium sulfate, to a final concentration of 1.2 M, was added to the concentrated A2058 conditioned medium for 1 hr. at 4°C. The solution was spun in a RC2-B Ultraspeed Sorvall centrifuge at 10,000 x g for 15 min. Only the supernatant had the capacity to stimulate motility.

In the first step, the sample was fractionated by hydrophobic interaction chromatography using 200 ml phenyl Sepharose CL-4B column equilibrated into 50 mM Tris (pH 7.5), 5% (v/v) methanol and 1.2 M ammonium sulfate. The supernatant from the ammonium sulfate fractionation was added to this column and eluted using linear gradients of 50 mM Tris (pH 7.5), 5% (v/v) methanol, with decreasing (1.2-0.0) M ammonium sulfate and increasing (0.50) % (v/v) ethylene glycol at 1 ml/min.

The active peak was pooled, dialyzed into 50 mM Tris, 0.1 M NaCl, 0.01 M CaCl<sub>2</sub>, 20% (v/v) ethylene glycol, and subjected to a second fractionation by lectin affinity chromatography using a 40 ml Affi-Gel concanavalin A column run at 1 ml/min. The sample was eluted in a stepwise fashion in the same buffer with 0, 10, and 500 mM methyl  $\alpha$ -mannopyranoside added successively. Fractions from each step of the gradient were pooled and tested for their capacity to stimulate motility.

In the third purification step, the sample that eluted at 500 mM  $\alpha$ -methyl-mannopyranoside was dialyzed into 10 mM Tris (pH 7.5) with 30% (v/v) ethylene glycol and fractionated by weak anion exchange chromatography. Chromatography was carried out on a ZORBAX BioSeries-WAX column using a Shimadzu BioLiquid chromatograph and eluted with a linear gradient of (0.0 - 0.4 M) sodium chloride at 3 ml/min.

The active peak was pooled, dialyzed against 0.1 M sodium phosphate (pH 7.2), 10% (v/v) methanol, and 10%

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(v/v) ethylene glycol, and subjected to a fourth fractionation step on a series of Spherigel TSK columns (4000SW, 4000SW, 3000SW, 2000SW, in that order). This molecular sieve step was run using the Shimadzu BioLiquid chromatograph at 0.4 ml/min.

The active peak was pooled and dialyzed into 10 mM Tris (pH 7.5), 5% (v/v) methanol, 20% (v/v) ethylene glycol and subjected to a fifth (strong anion exchange) chromatography step, a Pro-Pac PA1 column run at 1 ml/min using a Dionex BioLC with AI450 software. The sample was eluted with a linear gradient of (0.0-0.4M) NaCl.

In order to calculate activity yields after each step of purification, a unit of activity had to be derived. The dilution curve of ATX was biphasic with a broad peak and a linear range at sub-optimal concentrations. One unit of activity/well (i.e., 40 units/ml) was defined as 50% of the maximal activity in a full dilution curve. allowed calculation of the activity contained in any volume from the dilution needed to achieve 1 unit/well. Therefore, if a 1:10 dilution were needed in order to produce 1 unit of activity/well, the material contained 10 x 40 = 400 units/ml.Gel Electrophoresis. Protein samples were analyzed by SDSpolyacrylamide gel electrophoresis using the conditions of Laemmli (Laemmli, 1970). In brief, 7 or 8% SDS-containing polyacrylamide gels were prepared or pre-poured (8-16%) gradient gels were obtained commercially. Samples were prepared with or without reducing conditions (5%  $\beta$ -

gels were stained using Coomassie Blue G-250 as previously described (Neuhoff, et al., 1988). In this staining protocol, which ordinarily requires no destaining step, the Coomassie stain appears to be able to stain as little as 10 ng of protein.

mercaptoethanol). After electrophoretic separation, the

For two-dimensional electrophoresis, the protein, in 20% ethylene glycol, was dried in a Speed-vac and

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- redissolved in loading solution: 9M urea, 1% (v/v) pH 3-10 Bio-Lyte, and 2.5% (v/v) Nonidet-P40. This sample was then subjected to isoelectric focusing (O'Farrell, 1975) using a Bio-Rad tube cell in 120 x 3 mm polyacrylamide tube gels containing 9M urea, 2% (v/v) pH 3-10 Bio-Lyte,
- 0.25% (v/v) pH 8-10 Bio-Lyte and 2.5% (v/v) Nonidet-P40. 5 Reservoir solutions were 0.01 M phosphoric acid and 0.02 M NaOH. Non-equilibrium isoelectric focusing (O'Farrell, et al., 1977) was run initially with constant voltage (500 v) for 5 hr. Since the protein was basic, the procedure was
- repeated under equilibrium conditions (500 v for 17 hr.). 10 Electrophoresis in the second dimension was performed on a 7.5% polyacrylamide using the conditions of Laemmli The gel was stained with Coomassie Blue G-250 as (1970). above.
- Preparation of peptides for internal sequence of 15 autotaxin. Homogeneous ATX was sequentially digested with cyanogen bromide and, following reduction and pyridylethylation, with trypsin (Stone, et al., 1989). The resulting fragments were then separated by gradient
- elution on an Aquapore RP300 C-8 reverse phase column: 20 (v/v) trifluoroacetic acid and (0-70)% acetonitrile over 85 min. at a flow rate of 0.2 ml/min. A Dionex AI450 BioLC system was utilized and fractions were collected manually while monitoring the absorbance at 215 nm.
- Sequence analysis of peptides. The amino acid sequences 25 of peptides resulting from digestion and purification of ATX peptides #1-7 and 12-18, corresponding to SEQ ID NO:1 through SEQ ID NO:7 and SEQ ID NO:26 through SEQ ID NO:32, respectively, were determined on a Porton Instruments 2020 off-line sequenator using standard program #1.
- 30 Phenylthiohydantoin amino acid analysis of sequenator runs were performed on a Beckman System Gold HPLC using a modified sodium acetate gradient program and a Hewlett-Packard C-18 column. ATX-100 (SEQ ID NO:8), ATX-101 (SEQ ID NO:9), ATX-102 (SEQ ID NO:10), ATX-103 (SEQ ID NO:11)

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and ATX 104 (SEQ ID NO:33) were sequenced from gel-purified ATX.

Protein databases (Pearson, et al. 1988) that were searched for homologies in amino acid sequence with the ATX peptides include: GenBank (68.0), EMBL (27.0), SWISS-PROT (18.0), and GenPept (64.3).

#### EXAMPLE 1

## Purification of Autotaxin

The A2058 cells had been previously shown to produce protein factors which stimulate motility in an autocrine fashion (Liotta, et al., 1986). Conditioned medium from these cells was therefore used to identify and purify a new motility-stimulating factor, which is here named autotaxin and referred to as ATX. Since the purification was monitored with a biological assay, motility-stimulating activity had to be maintained throughout. The activity proved to be labile to freezing, acidic buffers, proteases (but not DNase or RNase), reduction, strong chaotrophic agents (e.g. > 4 M urea), and a variety of organic solvents (isopropanol, ethanol, acetonitrile). An organic solvent, ethylene glycol, which did not decrease bioactivity, was added for both storage and chromatographic separation.

100-200 L of serum-free conditioned medium were required in order to produce enough ATX for amino acid sequence analysis. The medium contained low concentrations of both BSA (5  $\mu$ g/ml) which was needed as a carrier protein and insulin (10  $\mu$ g/ml) which was required to support cell growth in low protein medium. Ultrafiltration to concentrate this large volume was performed with low protein-binding YM30 membranes which retain molecular species with M, > 30,000. As seen in Table 1, 200 L of conditioned medium prepared in this manner resulted in 10 x 10<sup>6</sup> units of activity. However, the initial unfractionated conditioned medium contained additional substances known to stimulate activity,

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particularly insulin, which does not completely wash out in the ultrafiltration step and which is additive to the motility stimulating activity in a complex manner (Stracke, et al., 1989). This had to be taken into account in order to determine yields for subsequent steps in which insulin had been removed.

TABLE 1. PURIFICATION OF AUTOTAXIN

10	Purification Step	Protein	Activity a	Specific	
	Recovery	(mg)	(total units)	Activity (units/mg)	(%) <sup>b</sup>
	200 L Conditioned Medium	33,000	10,000,000°	300	•
15	Phenyl Sepharose	1,235	460,000	370	100
	Concanavalin A	58	660,000	11,400	100
	Weak Anion Exchange	4.5	490,000	110,000	100
	TSK Molecular Sieves	~0.4 <sup>d</sup>	220,000	550,000	48
20	Strong Anion Exchange	~0.04 <sup>d</sup>	24,000 <sup>e</sup>	600,000	5.2

<sup>&</sup>lt;sup>a</sup> Activity calculated from Boyden chamber assay. The dilution which resulted in 50% of maximal activity (generally approximately 20 laser density units or ~40 cells/HPF) was chosen to have 1 unit of activity per well (equivalent to 40 units/ml).

- 25 b Recovery was estimated from activity, after the first purification column (i.e., phenyl sepharose).
  - <sup>e</sup> Initial activity in the unfractionated conditioned medium reflected the fact that insulin was used in the medium as a necessary growth factor under low protein conditions.
  - d Estimated protein is based on quantification by amino acid analysis.
- <sup>e</sup> This specific activity for purified protein corresponds to -10 fmol ATX/unit of motility activity (in a Boyden chamber well).

The first step in the purification involved fractionation by hydrophobic interaction chromatography using a phenyl Sepharose CL-4B column. The results are

shown in Figure 1. Most proteins, including insulin, eluted from the column in early fractions or in the void. However, the peak of activity eluted relatively late. The activity which was purified was estimated as 460,000 units ± 20% (Table 1). As the pooled peak of activity from the phenyl Sepharose fractionation is considered to be the first sample without significant insulin contamination, subsequent yields are measured against its total activity. Gel electrophoresis of a small portion of the pooled peak of activity (Figure 6A, column 2) revealed a large number of protein bands with BSA predominant from the original conditioned medium.

In the second step of purification, the active peak was applied to the lectin affinity column, Affi-Gel concanavalin A. As shown in Figure 2, most protein (estimated to be 90% of the total absorbance at 280 nm) failed to bind to the column at all. The non-binding fraction contained essentially no motility-stimulating activity (see dotted line in Figure 2). When a linear gradient of methyl  $\alpha$ -D-mannopyranoside was applied to the column, chemotactic activity eluted off in a prolonged zone, beginning at a concentration of approximately 20 mM sugar. Consequently, a step gradient was used to elute. Pure BSA failed to bind to con A.

Activity was found primarily in the 500 mM step of methyl  $\alpha$ -D-mannopyranoside. There appeared to be no significant loss of activity as seen in Table 1; however, specific activity (activity/mg total protein) increased thirty-fold. Gel electrophoresis of the pooled and concentrated peak (Figure 6A, column 3) revealed that the BSA overload was no longer apparent and the number of bands were much reduced. When the unbound protein was concentrated and applied to a gel, it appeared identical to the active peak from phenyl Sepharose-4B with a large BSA band.

The third purification step involved

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fractionating the previous active peak by weak anion exchange chromatography as shown in Figure 3. Under the running conditions, activity eluted in a broad peakshoulder or double peak in the middle of the shallow portion (0.0-0.4 M) of the NaCl gradient. The largest proportion of protein, lacking in motility-stimulating 5 capacity, bound strongly to the column and eluted off in high salt (1 M NaCl). There appeared to be no significant loss of activity, though specific activity increased by twenty-fold (Table 1). Analysis by gel electrophoresis of both the peak (28-34 min. in Figure 3) and the shoulder 10 (35-45 min. in Figure 3) is shown in Figure 6A (columns 4 and 5, respectively). Two predominant protein bands resulted: a broad doublet around 25-35 kDa and a second doublet around 110-130 kDa.

In the fourth purification step, the active peak was applied to a series of molecular sieves. Spectrophotometric monitoring of the eluant revealed two large peaks of protein (Figure 4). Activity corresponded to the first, higher molecular weight peak. Recovery of activity was ~48% with a five-fold increase in specific activity. Analysis by gel electrophoresis was performed under non-reducing and reducing conditions as shown in Figure 6B (columns 2 and 3, respectively). fractionation step had essentially removed all contaminating protein of molecular weight < 55 kDa. predominant band remaining has a molecular weight of 120 kDa unreduced and 125 kDa reduced; there are two minor bands with molecular weights 85 kDa and 60 kDa. that the 120 kDa protein changes so little in electrophoretic mobility after reduction would tend to indicate a paucity of disulfide bonds. However, the existing disulfide bonds have functional significance because motility-stimulating activity is labile to

reduction.

The fifth purification step involved

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fractionation of the active peak by strong anion exchange chromatography. As shown in Figure 5, activity corresponds to two broad optical absorbance peaks in the middle of the gradient with contaminating proteins eluting earlier. These two peaks were identical by amino acid analysis and by polyacrylamide gel electrophoretic separation. presumably represent different glycosylation states of the same parent protein. Activity is shown in Figure 5 at two different sample dilutions. Several dilutions of the fractionated samples were often necessary in order to resolve the true "peak" of activity as the shape of the ATX dilution curve was not sharp due to saturation and down regulation at high concentrations. Recovery from this chromatographic step is lower (5% compared to phenyl Sepharose), as might be expected when a minute quantity of protein is applied to a column; however, specific activity again increased (Table 1). Analysis by gel electrophoresis revealed a single protein band at molecular weight 120 kDa, unreduced (Figure 6C, column 2).

### EXAMPLE 2

20 <u>Characterization of Autotaxin</u>

Two dimensional gel electrophoresis of the purified protein (Figure 7) revealed a single predominant band. The band slopes downward slightly toward the basic side of the gel in a manner that is characteristic of glycosylated proteins. A basic pI of  $7.7 \pm 0.2$  was essentially the same whether the isoelectric focusing was run under non-equilibrium conditions (5 hr.) or was allowed to go to equilibrium (17 hr.).

A dilution curve of the purified protein is shown in Figure 8. The protein is active in the picomolar range and 1 unit of activity appears to correspond to a concentration of 400-600 picomolar (or approximately 10 fmol of ATX/Boyden chamber well). When dilutions were begun at higher concentrations of ATX, the resultant curve showed a broad plateau with down-regulation at the highest

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concentrations. The motility response to purified autotaxin is highly sensitive to pertussis toxin (hereinafter referred to as "PT") (Table 2 and Figure 9) with approximately 95% inhibition of activity at 0.5  $\mu$ g/ml PT.

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TABLE 2. Effect of Pertussis Toxin (PT) on Autotaxin-stimulated motility

A2058 Motility Response (density units <sup>1</sup> )			
control cells <sup>2</sup>	Pertussis 1	oxin-treated cells <sup>3</sup>	
Condition medium <sup>4</sup>	60.3	0.4	
Purified Autotaxin	38.5	0.0	

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Checkerboard analysis was performed to assess the random (chemokinetic) versus the directed 25 (chemotactic) nature of the motility response to ATX. Chambers were assembled with different concentrations of ATX above and below the filter, using ATX purified through the weak anion exchange fractionation step. Squares below the diagonal reflect response to a positive gradient, 30 squares above reflect response to a negative gradient, and squares on the diagonal reflect random motility in the absence of a gradient. ATX stimulates both chemotactic and chemokinetic responses (Figure 10), with chemotactic responses as high as fifteen-fold above background and 35

<sup>&</sup>lt;sup>1</sup> Chemotaxis quantitated by motility assay (Stracke, et al., 1978).

 $<sup>^2</sup>$  A2058 cell suspended at 2 x  $10^6$  cells/ml in DMEM supplemented with 1 mg/ml bovine serum and rocked at room temperature for 1 hr.

<sup>&</sup>lt;sup>3</sup> As control with 0.5  $\mu$ g/ml pertussis toxin.

<sup>&</sup>lt;sup>4</sup> Prepared by adding DMEM without phenol red supplemented with 0.1 mg/ml bovine serum albumin to subconfluent flasks of A2058 cells. The medium was harvested after 2 days incubation at 37°C in a humidified atmosphere and concentrated 25-30 fold using an Amicon ultrafiltration assembly with a YM-30 membrane.

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chemokinesis as high as eight-fold above background.

Amino acid analysis after complete acid
hydrolysis was used to quantitate purified protein. This
hydrolysis was carried out on protein excised from a
polyacrylamide gel and presumed to be pure. The analysis
indicated that 2.7 nmol of protein was present after
fractionation on the molecular sieve. After fractionation
by strong anion exchange chromatography, approximately 300
pmol remained. The results of the analysis are shown in
Table 3.

10 TABLE 3. AMINO ACID COMPOSITION OF AUTOTAXIN (CYS and TRP were not determined in this analysis)

	Amino Acid	Residues/100	
	ASX		12.5
	THR		6.0
	SER	•	5.7
15	GLX		9.4
	PRO		7.4
	GLY	•	7.0
	ALA		3.9
	VAL		6.7
	MET		1.2
	ILE		4.3
20	<b>LE</b> U		9.0
	TYR		5.2
	PHE		5.2
	HIS		3.8
	LYS		7.4
	ARG		5.4

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### EXAMPLE 3

# ATX Degradation and Determination of Amino Acid Sequence

information from purified ATX repeatedly proved futile.
The purified protein was therefore, sequentially digested and the resulting peptides fractionated by reverse phase chromatography. The results are shown in Figure 11.
Multiple sharp peaks including clusters at both the hydrophilic and hydrophobic ends of the gradient are seen.

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Several of these peptide peaks were chosen randomly for Edman degradation and N-terminal amino acid sequence analysis. Seven of the eight peaks (shown in Figure 11) chosen gave clear single sequence information as seen in Table 4. Using material from a separate digestion and purification, the remaining four sequences were also obtained.

Separate sense and antisense oligonucleotide probes were synthesized according to the fragment sequences of Table 4 by methods known to one skilled in the art. Representative probes are shown in Table 5.

TABLE 4. Peptide sequences for Autotaxin.

	PEPTIDE NO.	AMINO ACID SEQUENCE	SEQ ID: NO:	NAME
	1.	WHVA	SEQ ID NO:1	ATX 18
15	2.	PLDVYK	SEQ ID NO:2	ATX 19
	3.	YPAFK	SEQ ID NO:3	ATX 20
	4.	QAEVS	SEQ ID NO:4	ATX 24
	5.	PEEVTRPNYL	SEQ ID NO:5	ATX 29
20	6.	YDVPWNETI	SEQ ID NO:6	ATX 47
20	7.	VPPFENIELY	SEQ ID NO:7	ATX 48
	8.	GGQPLWITATK	SEQ ID NO:8	ATX 100
	9.	VNSMQTVFVGY-	SEQ ID NO:9	ATX 101
		GPTFK		
25	10.	DIEHLTSLDFFR	SEQ ID NO:10	ATX 102
	11.	TEFLSNYLTNVDD-	SEQ ID NO:11	ATX 103
		ITLVPETLGR		
	12.	QYLHQYGSS	SEQ ID NO:26	ATX 37
	13.	VLNYF	SEQ ID NO:27	ATX 39
30	14.	YLNAT	SEQ ID NO:28	ATX 40
	15.	HLLYGRPAVLY	SEQ ID NO:29	ATX 41
	16.	SYPEILTPADN	SEQ ID NO:30	ATX 44
	17.	XYGFLFPPYLSSSP	SEQ ID NO:31	ATX 53
35	18.	TFPNLYTFATGLY	SEQ ID NO:32	ATX 59

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# 19. VNVISGPIFDYDYDGLH SEQ ID NO:33 ATX 104 DTEDK

Peptide numbers 1-7 refer to peaks numbered in Figure 11. Peptide numbers 12-18 refer to peptides purified from the preparation which yielded peptide numbers 1-7. Peptides 8-11 and 19, are from a separate purification, not shown in Figure 11.

X refers to potentially glycosylated residues.

## TABLE 5.

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Oligonucleotides synthesized from peptide sequences of autotaxin (ATX).

The number of the oligonucleotide corresponds to the ATX peptide number as per Table 4. The final letter suffix distinguishes whether the oligonucleotide is a sense (S) or antisense (A) sequence.

	<u>Oligo</u>	Sequence	SEO ID NO:
15	A-18A	GTT-GGC-AGC-NAC-RTG-CCA	SEQ ID NO:12
	A-18S	TGG-CAY-GTN-GCT-GCC-AAC	SEQ ID NO:13
	A-20A	CTT-GAA-GGC-AGG-GTA	SEQ ID NO:14
	A-20S	TAY-CCT-GCN-TTY-AAG	SEQ ID NO:15
	A-29A	GGT-NAC-YTC-YTC-AGG	SEQ ID NO:16
20	A-29S	CCT-GAR-GAR-GTN-ACC	SEQ ID NO:17
	A-47A	NGT-NGC-RTC-RAA-TGG-CAC-RTC	SEQ ID NO:18
	A-47S	GAY-GTG-CCA-TTY-GAY-GCN-ACN	SEQ ID NO:19
	A-48A	GTT-DAT-RTT-STC-RAA-TGG-GGG	SEQ ID NO:20
	A-48S	CCC-CCA-TTT-GAG-AAC-ATC-AAC	SEQ ID NO:21
25	A-100A	CTT-NGT-NGC-NGT-DAT-CCA-NAR-	SEQ ID NO:22
		GGG-YTG-GCC-GCC	
	A-100S	GGC-GGC-CAR-CCC-YTN-TGG-ATH-	SEQ ID NO:23
		ACN-GCN-ACN-AAG	
30	A-101A	CTT-RAA-GGT-GGG-GCC-RTA-GCC-	SEQ ID NO:24
		CAC-RAA-GAC-TGT-YTG-CAT	
	A-101S	ATG-CAR-ACA-GTC-TTY-GTG-GGC-	SEQ ID NO:25
		TAY-GGC-CCC-ACC-TTY-AAR	

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## EXAMPLE 4

# Antipeptide Antibodies

Rabbits were injected with ATX-101 (SEQ ID NO:10) which had been cross-linked to bovine serum albumin. Antisera from these rabbits was subjected to salt precipitation followed by purification using affinity chromatography with Affi-Gel 10 beads covalently linked to the peptide, ATX-101 (SEQ ID NO:10). This affinity purified antibody reacted with the partially purified protein on immunoblots. This same antibody has been used to perform immunohistochemical stains on human tissue.

## EXAMPLE 5

# Enzymatic Deglycosylation of ATX

Purified ATX that was to be treated with peptide N-glycosidase F (PNGase F) was first dialyzed into 0.2 M sodium phosphate, 10% (v/v) ethylene glycol pH 7.0, using Centricon-30 ultrafiltration tubes. Varying concentrations of PNGase F were added to the ATX and incubated 16-18 hr. at 37°C. Complete digestion appeared to occur at concentrations of enzyme above 30 mU/ml (where 1 U converts 1mmol of substrate/min). For comparison, the experiments were repeated in the presence of 0.1 M  $\beta$ mercaptoethanol or 0.1% (w/v) SDS plus 0.5% (v.v) Nonidet-ATX that was to be treated with neuraminidase or 0glycosidase was dialyzed into 20 mM sodium phosphate, 0.1 M calcium acetate, and 10% (v/v) ethylene glycol (pH 7.2). Neuraminidase was added to a final concentration of 2 For treatment with neuraminidase alone, this mixture was incubated 16-18 hr at 37°C. Since Oglycosidase requires the removal of terminal sialic acid residues for efficient deglycosylation, ATX was preincubated with neuraminidase for 30-125 mU/ml and incubated 16-18 hr. at 37°C. The treated ATX was then dialyzed into 50 mM Tris with 20% ethylene glycol for storage at 5%C.

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Treatment of ATX with N-glycosylation altering agents A2058 cells were split into four 150 cm<sup>2</sup> flasks and incubated until just subconfluent in DMEM supplemented with 10% fetal calf serum. The medium was then replaced with fresh 10% FCS/DMEM to which had been added DPBS for control, 1mM dMAN, 1 mM NMdNM, or 10 mM (1.7 mg/ml) Swn. Concentrations of these pharmacological agents were similar to those previously described as inhibiting Nglycan processing enzymes in melanoma cells (Seftor, et al. 1991; Dennis, et al. 1990) as well as carcinoma cells (Ogier, et al. 1990). On the next day, each flask was washed twice with Dulbecco's phosphate buffered saline with calcium ("DPBS") then 20 ml of Dulbecco's minimum essential medium ("DMEM") supplemented with 0.01% (w/v)bovine serum albumin ("BSA") was added. The same concentration of each agent was added to the appropriate equilibrated flask and incubated for ~ 24 hr, after which the medium from each treatment group was collected, concentrated, washed into DPBS and stored at 5°C.

Cells from each flask were trypsinized and counted. There was no loss of viability or reduced cell number in any of the treatment groups compared to control cells.

## Effect of PNGase F on ATX

ATX binds to concanavalin A ("Con A") agarose beads and is eluted with buffer containing 0.5 M methyl a-D-mannopyranoside, indicating that ATX is likely to contain mannose residues. Such mannose sugar residues are most characteristic of N-linked oligosaccharides. In order to verify that ATX contained asparagine-linked oligosaccharides, we treated it with the endoglycosidase, PNGase F, which cleaves high mannose, hybrid, and complex N-linked oligosaccharides at the asparagine residue.

Partially purified ATX was treated with 60 mU/ml of enzyme under a variety of increasingly denaturing conditions and then separated by polyacrylamide gel

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electrophoresis (Figure 16). Lane 1 shows untreated material; the 125 kDA band (arrow) is autotaxin. this material is treated overnight with PNGase F under very mild conditions, the size of the 125 kDa band decreases to ~100- 105 kDa. Addition of 0.1 M bmercaptoethanol (Lane 2) or 0.5% Nonidet-P40 (lane 3) to 5 the ATX sample has no effect on the size of the resultant protein band. Even complete denaturation of ATX of boiling the sample for 3 min in 0.1% SDS with (lane 5) or without (lane 4)  $\beta$ -mercaptoethanol, followed by addition of 0.5% Nonidet-P40 to maintain enzymatic activity, has no 10 effect on the final size of deglycosylated protein,

indicating that the deglycosidation reaction was complete even under mild conditions.

Because these results showed that ATX contained N-linked oligosaccharide groups, it became important to 15 see if these sugar moieties were necessary for stimulation of motility. The partially purified ATX sample was treated with varying concentrations of PNGase F (0.1 to 60 mU/ml) under mild, non-denaturing conditions. Analysis of the resulting digest by polyacrylamide gel electrophoresis 20 is shown in Figure 17A. As this figure shows, the digestion was incomplete using from 0.1 to 10 mU/ml of enzyme and resulted in a smear of protein between 100-125 However, at higher concentrations of enzyme, cleavage of N-linked oligosaccharides from ATX appears to 25 be complete. When these different digestion products were compared for their capacity to stimulate motility (Figure 17B), there was no significant difference between groups.

### EXAMPLE 6

Cloning the 3' end of Autotaxin (4C11) ATX is active in picomolar to nanomolar concentrations and is synthesized in very small concentrations by A2058 cells. As might be expected, the cDNA clone was relatively rare, requiring various strategies and multiple library screenings in order to

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identify it (Figure 12). Attempts to utilize degenerate oligonucleotides deduced from known peptide sequences were unsuccessful--whether we used the oligo nucleotides for screening cDNA libraries or for reverse transcription of mRNA followed by amplification with the polymerase chain reaction (RT/PCR). We then utilized an affinity-purified anti-peptide ATX-102 antibodies to screen an A2058 expression library.

These anti-peptide antibodies were generated by methods well established in the art and described 10 previously with slight modification (Wacher, et al., In brief, the previously identified peptide, ATX-102 (Stracke, et al., 1992), was synthesized on a Biosearch 9600 peptide synthesizer. It was then solubilized in 1X PBS containing 20% (v/v) DMSO and 15 conjugated to the protein carrier, bovine serum albumin (BSA), with glutaraldehyde. For the first injection into New Zealand white rabbits, the BSA-peptide conjugate was emulsified with complete Freund's adjuvant and injected subcutaneously. For subsequent injections, the BSA-20 peptide conjugate was emulsified with incomplete Freund's adjuvant. The resultant antiserum was heat-inactivated at 56°C for 30 min. Immunoglobulins were precipitated out in 47% saturated ammonium sulfate, then redissolved and dialyzed into PBS. Antibodies were adsorbed onto peptideconjugated Affi-Gel 10 resin (made using the BioRad 25 protocol), eluted with 0.1 N acetic acid, and neutralized with 2 M Tris-HCl, pH 8. The resulting affinity-purified antibodies were dialyzed into DPBS, concentrated, aliquotted, and stored at -20°C. The antibodies were found to recognize a 125 kDa protein on immunoblots of 30 partially purified A2058 conditioned medium and to preferentially stain some breast carcinoma cells compared to normal breast using immunohistochemical techniques.

An A2058 cDNA library was prepared by purifying poly-A purified mRNA from the cells then size-selecting

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• mRNA > 1000 bp for the preparation of cDNA. The cDNA inserts were placed into λgt11 directionally, using the ProMega cDNA kit using standard methods well-established in the field. LE 392 cells were infected with the λgt11 and plaques were transferred onto nitrocellulose membranes by overnight incubation at 37°C. The antibody was incubated with the membranes in blocking buffer for 2 hr at room temperature, using approximately twice the concentration of antibody which gave a strong response on Western blot analysis. Secondary antibody was goat anti-rabbit immunoglobulin, and the blot was developed colorimetrically with 4-chloro-1-naphthol.

Positive clones were confirmed by antibody competition with specific peptides but not unrelated peptides. Using this technique and multiple subclonings, we obtained a partial cDNA clone of the autotaxin gene, which we called 4C11. The 4C11 insert was removed from Agt11 by restriction enzyme digests and subcloned into pBluescript for sequencing by standard Sanger techniques (Sanger, et al., 1977). The 4C11 clone contained bases, including the poly-adenylated tail and the AATAAA adenylation signal locus, i.e., it contained the 3' terminus of the gene. It also included a 627 base open reading frame. Database analysis of this nucleotide sequence revealed that it is unique. The predicated amino acid sequence for 4C11 is 209 amino acids long with exact matches for 7 previously identified ATX peptides: (ATX-20, ATX-34, ATX-102, ATX-104, ATX-204, ATX-215, and ATX-244).

### EXAMPLE 7

# Cloning the 5' terminus of ATX

Database analysis of the 3' terminus of the ATX gene demonstrated a novel protein. However, we have found an interesting homology that has helped to guide us in exploring its function. ATX had a 45% amino acid identity and a 57% nucleotide identity with PC-1, a marker of B cell activation found on the surface of plasma cells.

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Using the PC-1 protein sequence as a guide, we found that ATX peptide homologies were scattered throughout the length of the protein. The only exception was the far amino terminus of PC-1, which includes the transmembrane and intracellular domains, and which had no homologies.

Knowing approximate localization of the ATX peptides along the length of ATX, we then amplified different segments of ATX by the PCR (Figure 13). These amplified segments of DNA were then subcloned into plasmids utilizing the TA Cloning kit of ProMega. The PCR amplified DNA could then be sequenced using standard Sanger sequencing techniques (Sanger, et al., 1977).

### Cloning of full length ATX gene

A reverse transcriptase reaction was performed using total or oligo-(dT) purified RNA from A2058 or N-tera 2D1 cells as template and an anti-sense primer from the 5' end of 4C11 (GCTCAGATAAGGAGGAAAGAG). This was followed by one or two PCR amplification of the resultant cDNA using the commercially available kit from Perkin-Elmer and following manufacturer's directions.

These PCR reactions utilized nested antisense primers from 4C11 (GAATCCGTAGGACATCTGCTT and TGTAGGCCAAACAGTTCTGAC) as well as degenerate, nested sense primers deduced from ATX peptides: ATX-101 (AAYTCIATGCARACIGTITTYGTIG and TTYGTIGGITAYGGICCIACITTYAA), ATX-103

25 (AAYTAYCTIACIAAYGTIGAYGAYAT and
GAYGAYATIACICTIGTICCIGGIAC), or ATX-224
(TGYTTYGARYTICARGARGCIGGICCICC). The amplified DNA was
then purified from a polyacrylamide gel using standard
procedures and ligated into the pCR™ plasmid using the TA
30 cloning kit (Invitrogen Corporation) according to
manufacturer's directions.

The 5' RACE kit was utilized to extend the 5' end of ATX cDNA using total RNA from N-tera 2D1 as template and previously obtained sequence as primer (GCTGTCTTCAAACACACGC). The 5' end of the A2058 synthesized

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protein was obtained by using previously obtained sequence as primer (CTGGTGGCTGTAATCCATAGC) in a reverse transcriptase reaction with total A2058 RNA as template, followed by PCR amplification utilizing the 5' end of N-tera 2D1 sequence as sense primer

5 (CGTGAAGGCAAAGAGAACACG) and a nested antisense primer (GCTGTCTTCAAACACAGC). A2058 DNA encoding ATX is set forth in a SEQ ID NO:68 and the amino acid sequence is provided in SEQ ID NO:69.

DNA sequencing: DNA sequencing was performed using dideoxy methodology (Sanger, et al. 1977) and (35S) dATP (Du Pont, New England Nuclear).

We have found one region between the 5' end of the 4Cl1 and the ATX peptide designated ATX-101, also referred to as the "hot spot". This region has been sequenced five times with different sequences found each time. The hot spot appears to be located within the region from approximately nucleotide 1670 to 1815. The consensus sequence is represented by amino acids position 559 through 604. Variations found include DNA sequence that results in single and multiple amino acid insertions. One sequence had a stop codon in this region and may have represented an intron. This region has been found to be variable in forms of ATX.

### EXAMPLE 8

25 Cloning ATX in a human teratocarcinoma cell line

The fact that ATX is present in other cancer
cells was confirmed by sequence information from N-tera
2D1, a human teratocarcinoma cell line. For these cells,
a prepared cDNA library in \(\lambda\text{gt10}\) was amplified and the

CDNA inserts were extracted. Using oligonucleotide
primers based on known A2058 ATX sequence, DNA segments
were amplified by PCR. The DNA segments were then
subcloned into plasmids and sequenced as for A2058. We
have 3104 bp DNA sequence for N-tera ATX (SEQ ID NO:66)
and smaller portions thereof. This includes an open

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reading frame that codes for a putative protein containing 861 amino acids (SEQ ID NO:67) and smaller portions thereof. Like the A2058 ATX, the N-tera 2D1 sequence has homologies for multiple ATX peptides (Figure 15). Sequence homology between the A2058 and N-tera 2D1 cells is approximately 99%.

#### EXAMPLE 9

Cloning 5' end of ATX in human normal liver
The 5' end of ATX has proven difficult to obtain

from either tumor cell line to date. Normal human liver mRNA was therefore amplified using the 5' RACE kit (Clontech) with known sequence from A2058 ATX as antisense primer. A DNA segment was obtained and has been sequenced. This segment codes for 979 amino acids, including an initiating methionine (SEQ ID NO:38). The putative protein sequence also includes a 20 amino acid transmembrane domain which is different from the tumor ATX's (SEQ ID NO:54), as shown in Table 7. Both tumorous forms of ATX apparently lack a transmembrane region and are instead secreted proteins.

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#### Table 7

Nucleotide and Amino Acid Sequences Encoding Liver ATX Amino Terminus containing the Transmembrane region

- 25 Protein Sequence (SEQ ID NO: 54)
  - Met Ala Arg Arg Ser Ser Phe Gln Ser Cys Gln Asp Ile Ser Leu Phe Thr Phe Ala Val Gly Val Asn Ile Cys Leu Gly Phe Thr Ala His Arg Ile Lys Arg Ala Glu Gly Trp
  - DNA Sequence (SEQ ID NO: 53)
- ATGGCAAGGA GGAGCTCGTT CCAGTCGTGT CAAGATATAT CCCTGTTCAC
  TTTTGCCGTT GGAGTCAATA TCTGCTTAGG ATTCACTGCA CATCGAATTA
  AGAGAGCAGA AGGATGG

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#### EXAMPLE 10

### Domains of ATX

Searches of protein databases (Pearson, et. al. 1988) confirmed that the homology between ATX and PC-1 was present throughout the length of the extracellular portion of the molecules (Buckley, et. al., 1990; Funakoshi, et. al. 1992). There is a 45% amino acid identity and a 64% similarity between the 2 protein sequences (Fig. 18). For the cDNA sequence, the identity is ~57%.

These proteins share several interesting properties and domains (Fig. 19). Both have a number of 10 potential N-linked glycosylation sites: four for ATX (Asn54, Asn463, Asn577, Asn859) and nine for PC-1. have adjacent somatomedin B domains near the amino end of the extracellular domain. This somatomedin B domain is a cysteine-rich region containing 3 presumed cystine cross-15 linkages. ATX has 33 Cys residues and PC-1 has 37; 30 of these Cys residues are identical in placement. proteins also contain the loop region of an EF hand (Buckley, et. al. 1990; Kretsinger, 1987). In addition, both proteins have a transmembrane/signal peptide region 20 with a short intracellular peptide, common in ectoenzymes (Maroux, 1987). However, the amino acid identity between ATX and PC-1 in the intracellular and transmembrane regions is only 11%.

Finally, both proteins have a region homologous to the bovine intestinal phosphodiesterase enzymatic domain with conversation of the threonine that is thought to act as the intermediate phosphate binding site (Culp, et al. 1985). PC-1 has been demonstrated to have phosphodiesterase type I, nucleotide pyrophosphatase, and threonine-specific kinase enzymatic activities (Rebbe, et al. 1991; Oda, et al. 1991). In order to test whether purified ATX had type I phosphodiesterase activity, samples were incubated with p-nitrophenyl thymidine-5'-monophosphate at pH 8.9 for 30 min. Samples were assayed

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- in a 100 µl volume containing 50 mM Tris-HCl, pH 8.9 and 5 mM p-nitrophenyl thymidine-5'-monophosphate. After a 30 minute incubation at 37 °C the reactions were terminated by addition of 900 ml 0.1 N NaOH and the amount of product formed was determined by reading the absorbance at 410 nm.
- ATX was found to hydrolyze the p-nitrophenyl thymidine-5'monophosphate (Razzell, 1963) at a rate of 10 pmol/ng/min,
  a reaction rate similar to that reported for PC-1 (Oda, et
  al. 1993).

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All publications mentioned hereinabove are hereby incorporated in their entirety by reference.

While the foregoing invention has been described in some detail for purposes of clarity and understanding, it will be appreciated by one skilled in the art from a reading of this disclosure that various changes in form and detail can be made without departing from the true scope of the present invention and appended claims.

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## SEQUENCE LISTING

•	•	SEQUENCE DISTING
	(1) GENE	RAL INFORMATION:
	(i)	APPLICANT: STRACKE, MARY; LIOTTA, LANCE; SCHIFFMANN, ELLIOTT; KRUTZSCH, HENRY; MURATA, JUN
5	(ii)	TITLE OF INVENTION: MOTILITY STIMULATING PROTEIN USEFUL IN CANCER DIAGNOSIS AND THERAPY
	(iii)	NUMBER OF SEQUENCES: 69
10	(iv)	CORRESPONDENCE ADDRESS:  (A) ADDRESSEE: MORGAN & FINNEGAN  (B) STREET: 345 PARK AVENUE  (C) CITY: NEW YORK  (D) STATE: NEW YORK  (E) COUNTRY: U.S.A.  (F) ZIP: 10154
15	(v)	COMPUTER READABLE FORM:  (A) MEDIUM TYPE: Floppy Disk  (B) COMPUTER: IBM PC compatible  (C) OPERATING SYSTEM: PC-DOS/MS-DOS  (D) SOFTWARE: WordPerfect 5.1
20	(vi)	CURRENT APPLICATION DATA: (A) APPLICATION NUMBER: (B) FILING DATE: 24-MAY-1995 (C) CLASSIFICATION:
	(vii)	PRIOR APPLICATION DATA (A) APPLICATION NUMBER: 08/346,455 (B) FILING DATE: 28-NOV-1994
25	(vii)	PRIOR APPLICATION DATA (A) APPLICATION NUMBER: 08/249,182 (B) FILING DATE: 25-MAY-1994
	(vii)	PRIOR APPLICATION DATA (A) APPLICATION NUMBER: 07/822,043 (B) FILING DATE: 17-JAN-1992
30	(viii)	ATTORNEY/AGENT INFORMATION: (A) NAME: DOROTHY R. AUTH (B) REGISTRATION NUMBER: 36,434 (C) DOCKET NUMBER: 2026-4149US2
35	(ix)	TELECOMMUNICATION INFORMATION: (A) TELEPHONE: (212) 758-4800 (B) TELEFAX: (212) 751-6849

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                      TYPE: amino acid
                 (D)
                      TOPOLOGY: linear
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                      LENGTH: 5 amino acids
                 (B)
                      TYPE: amino acid
                      TOPOLOGY: linear
                 (D)
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      (2)
           INFORMATION FOR SEQ ID NO:5:
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- 49 -

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	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 10</li><li>(B) TYPE: amino acid</li><li>(D) TOPOLOGY: linear</li></ul>
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	(2) INFORMATION FOR SEQ ID NO:8:
25	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 11</li> <li>(B) TYPE: amino acid</li> <li>(D) TOPOLOGY: linear</li> </ul>
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:8
30	Gly Gly Gln Pro Leu Trp Ile Thr Ala Thr Lys 1 5 10
	(2) INFORMATION FOR SEQ ID NO:9:
25	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 16</li></ul>

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(B) TYPE: amino acid

(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

Val Asn Ser Met Gln Thr Val Phe Val Gly Tyr Gly
1 5 10

5 Pro Thr Phe Lys

10

20

- (2) INFORMATION FOR SEQ ID NO:10:
  - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 12

(B) TYPE: amino acid

(D) TOPOLOGY: linear

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:
- Asp Ile Glu His Leu Thr Ser Leu Asp Phe Phe Arg

  15 1 0 10
  - (2) INFORMATION FOR SEQ ID NO:11:
    - (i) SEQUENCE CHARACTERISTICS:
      - (A) LENGTH: 23
      - (B) TYPE: amino acid
      - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:

Thr Glu Phe Leu Ser Asn Tyr Leu Thr Asn Val Asp 1 5 10

- 25 Asp Ile Thr Leu Val Pro Glu Thr Leu Gly Arg
  - (2) INFORMATION FOR SEQ ID NO:12:
    - (i) SEQUENCE CHARACTERISTICS:
      - (A) LENGTH: 18
- 30 (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:

.

35 GTTGGCAGCN ACRTGCCA

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0										
	(2)	INFO	RMATION FOR SEQ ID NO:13:							
5		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 18  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear							
		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:13:	·						
	TGGC	CAYGTN	IG CTGCCAAC	18						
10	(2) INFORMATION FOR SEQ ID NO:14:									
10		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 15  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear							
15		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:14:							
	CTT	GAAGGC	CA GGGTA	15						
	(2)	INFO	ORMATION FOR SEQ ID NO:15:							
20		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 15  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear							
		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:15:							
25	TAY	CCTGCI	'NT TYAAG	15						
	(2)	INF	ORMATION FOR SEQ ID NO:16:	·						
30		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 15  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear							
		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:16:							
25	GG"	INACYT	TCY TCAGG	15						

BNSDOCID: <WO\_\_\_9532221A2\_I\_>

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	(2)	INFO	RMATION FOR SEQ ID NO:17:	
5		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 15  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:17:	
	CCTG	SARGAR	G TNACC	15
10	(2)	INFO	RMATION FOR SEQ ID NO:18:	
15		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 21  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
•		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:18:	
	NGTI	NGCRTC	R AATGGCACRT C	21
20	(2)	INFO	RMATION FOR SEQ ID NO:19:	
20		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 21  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
25		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:19:	
	GAY	GTGCCA	T TYGAYGCNAC N	21
	(2)	INFO	RMATION FOR SEQ ID NO:20:	
30		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 21  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:20:	
35	GTT	DATRTI	S TCRAATGGGG G	21

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	(2)	INFO	RMATION FOR SEQ ID NO:21:	
5		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 21  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:21:	
	CCCC	CATTT	G AGAACATCAA C	21
10	(2)	INFO	DRMATION FOR SEQ ID NO:22:	
		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 33  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
15		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:22:	-
	CTT	NGTNG(	CN GTDATCCANA RGGGYTGGCC GCC	33
	(2)	INF	ORMATION FOR SEQ ID NO:23:	
20		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 33  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
25		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:23:	
	GGC	:GGCCA	RC CCYTNTGGAT HACNGCNACN AAG	33
	(2)	INF	FORMATION FOR SEQ ID NO:24:	
30		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 39  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	

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•	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:	
	CTTRAAGGTG GGGCCRTAGC CCACRAAGAC TGTYTGCAT	39
	(2) INFORMATION FOR SEQ ID NO:25:	
5	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 39  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
10	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:25:	
10	ATGCARACAG TCTTYGTGGG CTAYGGCCCC ACCTTYAAR	39
	(2) INFORMATION FOR SEQ ID NO:26:	
15	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 9</li> <li>(B) TYPE: amino acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:26:	
20	Gln Tyr Leu His Gln Tyr Gly Ser Ser 1 5	
	(2) INFORMATION FOR SEQ ID NO:27:	
25	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 5</li> <li>(B) TYPE: amino acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
•	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:	
30	Val Leu Asn Tyr Phe 1 5	
	(2) INFORMATION FOR SEQ ID NO:28:	
35	(i) SEQUENCE CHARACTERISTICS:	

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(A) LENGTH: 5

(B) TYPE: amino acid

- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear
- 5 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:

Tyr Leu Asn Ala Thr

- (2) INFORMATION FOR SEQ ID NO:29:
- 10 (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 11
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- 15 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:29:

  His Leu Leu Tyr Gly Arg Pro Ala Val Leu Tyr

  1 5 10
  - (2) INFORMATION FOR SEQ ID NO:30:
- 20 (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 11
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- 25 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:

  Ser Tyr Pro Glu Ile Leu Thr Pro Ala Asp Asn
  1 5 10
  - (2) INFORMATION FOR SEQ ID NO:31:

30 (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 14
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear

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0	•	(xi)	SE	QUE1	ICE I	DESCI	RIPT	ON:	SEQ	ID 1	10:3	l:
	Xaa 1 Ser	Tyr	Gly	Phe	Leu 5	Phe	Pro	Pro	Tyr	Leu 10	Ser	Ser
	ser	PIO										
5	(2)	INF	ORMA	TIOI	v FO	R SE	Q ID	NO:	32:			
		(i)	(A (E (C	(a) 1 (b) 5 (c) 5	LENG' LYPE STRAI	TH: : : am NDED	ACTEI 13 ino a NESS : lii	acid	•			
10		(vi)	C E	ירוביו	יזרב י	חדפת	ייים דם	TON:	SEQ	ו מד	NT() • 3 '	<b>9</b> •
	The	Phe										
	1 Tyr	rne	PIO	ASII	5	1y1	TIII	FIIC	AIA	10	GIY	Беа
15	(2)	INF	ORMA	TIO	N FO	R SE	QĮD	NO:	33:			
20		(i)	( <i>I</i> (E	4) : 3) : C) :	LENG TYPE STRA	TH: : am NDED	ino .	acid : si	ngle			
		(xi)	SI	EQUE	NCE	DESC	RIPT	ION:	SEQ	ID	NO:3	3:
	Val 1	Asn	Val	Ile	Ser 5	-	Pro	Ile	Asp	Asp	_	Asp
25	_	Asp	Gly 15	Leu	_		Thr	Glu 20		Lys		
	(2)	INE	FORM	ATIO	N FO	R SE	Q ID	NO:	34:			
30		(i)		(A (B (C	) I ) T	ENGT YPE:	H: am DEDN	829 ino ESS:	ISTI acid si know	ngle		
	•	·(ii	L)	МО	LECU	LE I	YPE:	pr	otei	n		
		(ii	ii)	НУ	POTH	ETIC	'AL:	No				
25		(vi	L)	OR	IGIN	IAL S	OURC	E:				

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ORGANISM: Human
                   (A)
                   (B)
                        STRAIN:
                        INDIVIDUAL ISOLATE:
                   (C)
                        DEVELOPMENTAL STAGE:
                   (D)
                        HAPLOTYPE:
                   (E)
                   (F)
                        TISSUE TYPE:
                        CELL TYPE: Melanoma
                   (G)
5
                   (H)
                        CELL LINE: A2058
                   (I)
                        ORGANELLE:
          (ix)
                  FEATURE:
                   (A)
                        NAME/KEY:
                   (B)
                        LOCATION:
                        IDENTIFICATION METHOD:
                   (C)
10
                        OTHER INFORMATION: Putative protein
                   (D)
                        sequence of A2058 Autotaxin
                   SEQUENCE DESCRIPTION: SEQ ID NO:34:
           (xi)
     Cys His Asp Phe Asp Glu Leu Cys Leu Lys Thr Ala
     Arg Gly Trp Glu Cys Thr Lys Asp Arg Cys Gly Glu
15
     Val Arg Asn Glu Glu Asn Ala Cys His Cys Ser Glu
     Asp Cys Leu Ala Arg Gly Asp Cys Cys Thr Asn Tyr
     Gln Val Val Cys Lys Gly Glu Ser His Trp Val Asp
     Asp Asp Cys Glu Glu Ile Lys Ala Ala Glu Cys Pro
20
                       65
     Ala Gly Phe Val Arg Pro Pro Leu Ile Ile Phe Ser
                                    80
               75
     Val Asp Gly Phe Arg Ala Ser Tyr Met Lys Lys Gly
                            90
     Ser Lys Val Met Pro Asn Ile Glu Lys Leu Arg Ser
                                       105
                  100
25
     Cys Gly Thr His Ser Pro Tyr Met Arg Pro Val Tyr
          110
                               115
     Pro Thr Lys Thr Phe Pro Asn Leu Tyr Thr Leu Ala
     Thr Gly Leu Tyr Pro Glu Ser His Gly Ile Val Gly
              135
     Asn Ser Met Tyr Asp Pro Val Phe Asp Ala Thr Phe
30
                           150
     His Leu Arg Gly Arg Glu Lys Phe Asn His Arg Trp
                  160
                                       165
      Trp Gly Gly Gln Pro Leu Trp Ile Thr Ala Thr Lys
      Gln Gly Val Lys Ala Gly Thr Phe Phe Trp Ser Val
                       185
```

•			Pro 195	•				200				
	205		Leu			210					215	
	Val	_	Ala	220					225			
5		230	Lys				235					240
3			Gly		245					250		
	_		Lys 255					260				
	265		Ala			270					275	
10		_	Met	280					285			
		290	Met				295					300
			Gly		305					310		
			Arg 315				•	320				
15	325		Gly			330					335	
			Leu	340					345			
		350	Leu Phe				355					360
20		-	Ile		365					370		
20			375 Phe					380				
	385		Leu			390					395	
	_		His	400					405			
25		410					415	5				420
			_		425					430		Asp
	_	-	435					440	)			. Gly
	445	_				450	)				455	Pro
30				460	)				465	5		Cys
		470	1				475	5				480 Gly
					485	,		s Lei	ı Let	490	)	Asn
25			495	,				500	)		Thi	r Arg
35	505					510					519	5

=												
	Pro	Asn	Tyr	Pro 520	Gly	Ile	Met	Tyr	Leu 525	Gln	Ser	Asp
	Asp	Asp 530	Leu		Cys	Thr	Cys 535	Asp		Lys	Val	Glu 540
	Pro	Lys	Asn	Lys	Leu 545	Asp		Leu	Asn	Lys 550	Arg	Leu
5	His	Thr	Lys 555	Gly		Thr	Glu	Glu 560	Arg	His	Leu	Leu
	Tyr 565	Gly	Arg	Pro	Ala	Val 570	Leu	Tyr	Arg	Thr	<b>Arg</b> 575	Tyr
	Asp	Ile	Leu	Tyr 580	His	Thr	Asp	Phe	Glu 585	Ser	Gly	Tyr
		Glu 590					595					600
10		Val		_	605					610		
	_	His	615			_		620				
	625	Ser				630					635	
15		Asn		640					645			
15		Pro 650	_				655					660
	_	Ala			665					670		
		Ala	675					680				
20	685	Leu				690					695	
		Asn		700		_			705			
		Asp 710					715					720
		Tyr			725					730		
25		Tyr	735					740				
	745	Gln Ser				750					755	
		Glu		760					765			
		770		_			775					780
30		Val			785					790		
		Arg	795					800				
	805	Arg	ьys	inr	ser	810		ıyr	F10	GIU	815	
35	Thr	Leu	Lys	Thr 820		Leu	His	Thr	Tyr 825		Ser	Glu

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	(2) INFORM	ATION FOR SEQ ID NO:35:	
5	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 2946  (B) TYPE: nucleic acid  (C) STRANDEDNESS: double  (D) TOPOLOGY: Unknown	
	(ii)	MOLECULE TYPE: cDNA	÷
10	(iii)	HYPOTHETICAL: No	
	(vi)	ORIGINAL SOURCE: (A) ORGANISM: Human (B) STRAIN: (C) INDIVIDUAL ISOLATE: (D) DEVELOPMENTAL STAGE:	
15		(E) HAPLOTYPE: (F) TISSUE TYPE: (G) CELL TYPE: Melanoma (H) CELL LINE: A2058 (I) ORGANELLE:	
20	(ix)	FEATURE: (A) NAME/KEY: (B) LOCATION: (C) IDENTIFICATION METHOD: (D) OTHER INFORMATION: Partial DNA Sequence of A2058 Autotaxin	e
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:35:	
25	TGGCTGGGAG AATGAAGAAA CCAGGGGAGA	TGTACTAAGG ACAGATGTGG AGAAGTCAGA ATGCCTGTCA CTGCTCAGAG GACTGCTTGG  CTGCTGTACC AATTACCAAG TGGTTTGCAA  1	40 80 20 60
	AAGGCCGCAG TAATCATCTT GAAGAAAGGC AGGTCTTGTG	AATGCCCTGC AGGGTTTGTT CGCCCTCCAT  CTCCGTGGAT GGCTTCCGTG CATCATACAT  AGCAAAGTCA TGCCTAATAT TGAAAAACTA  GCACACACTC TCCCTACATG AGGCCGGTGT  3	240 280 320 360
30	TGGGCTATAT ATGTATGATC GGCGAGAGAA GCTATGGATT	CCAGAATCAC ATGGAATTGT TGGCAATTCA CTGTATTTGA TGCCACTTTT CATCTGCGAG ATTTAATCAT AGATGGTGGG GAGGTCAACC ACAGCCACCA AGCAAGGGGT GAAAGCTGGA	140 180 520 560
35	TATTAACCAT GAGGCCTTCG	ATTGCGGTGG CTCACCCTGC CAGATCATGA GTCTATGCCT TCTATTCTGA GCAACCTGAT	640 680 720

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•					
	GTAGTTATGG	CTCACCTTTT	ACTCCGGCTA	AGAGACCTAA	760
	GAGGAAAGTT	GCCCCTAAGA	GGAGACAGGA	AAGACCAGTT	800
	GCTCCTCCAA	AGAAAGAAG	AAGAAAAATA	CATAGGATGG	840
	ATCATTATGC	TGCGGAAACT	CGTCAGGACA	AAATGACAAA	880
	TCCTCTGAGG	GAAATCGACA	AAATTGTGGG	GCAATTAATG	920
	CATCCACTCA	AACAACTAAA	ACTGCGTCGG	TGTGTCAACG	960
_	TO TOTAL	CGGAGACCAT	GGAATGGAAG	ATGTCACATG	1000
5	TCATAGAACT	GAGTTCTTGA	GTAATTACCT	AACTAATGTG	1040
	САТСАТАТТА	CTTTAGTGCC	TGGAACTCTA	GGAAGAATTC	1080
	CATCCAAATT	TAGCAACAAT	GCTAAATATG	ACCCCAAAGC	1120
	CATTATTGCC	AATCTCACGT	GTAAAAAACC	AGATCAGCAC	1160
	TTTAAGCCTT	<b>ACTTGAAACA</b>	GCACCTTCCC	AAACGTTTGC	1200
	ACTATGCCAA	CAACAGAAGA	ATTGAGGATA	TCCATTTATT	1240
	CCTCCAACGC	<b>AGATGGCATG</b>	TTGCAAGGAA	ACCTTTGGAT	1280
10	CTTTATAAGA	AACCATCAGG	AAAATGCTTT	TTCCAGGGAG	1320
	<b>ACCACGGATT</b>	TGATAACAAG	GTCAACAGCA	TGCAGACTGT	1360
	TTTTGTAGGT	TATGGCCCAA	CATTTAAGTA	CAAGACTAAA	1400
	GTGCCTCCAT	TTGAAAACAT		AATGTTATGT	1440
	GTGATCTCCT	GGGATTGAAG	CCAGCTCCTA	ATAATGGGAC	1480
	CCATGGAAGT	TTGAATCATC	TCCTGCGCAC	TAATACCTTC	1520 1560
	AGGCCAACCA	TGCCAGAGGA	AGTTACCAGA	CCCAATTATC	1600
15	CAGGGATTAT	GTACCTTCAG	TCTGATTTTG	ACCTGGGCTG	1640
13	CACTTGTGAT	GATAAGGTAG	AGCCAAAGAA	CAAGTTGGAT	1680
	GAACTCAACA	AACGGCTTCA	TACAAAAGGG	TCTACAGAAG	1720
	AGAGACACCT	CCTCTATGGG	CGACCTGCAG	TGCTTTATCG	1760
	GACTAGATAT	GATATCTTAT	ATCACACTGA	CTTTGAAAGT	1800
	GGTTATAGTG	AAATATTCCT	AATGCIACIC	TGGACATCAT	1840
	ATACTGTTTC	CAAACAGGCT	CCCCTCATCT	GCGTTCCTGA CCGTGTTTCT	1880
20	CCATCTGACC	AGTIGCGICC	. GGCCIGAIGI · TTTCCCCCTA	AAAAATGATA	1920
20	ACCACATTICA	GICAGAACIC	, LIIGGCCIA(	CTTATCTGAG	1960
	AGCAGATGTC	CIACGGAIIC	· ATCATCCATT	CCTTGTAACC	2000
	A A TRA TROCTUTO	CNATCTATC	TCCTTTCAA	CGGGTCTGGA	2040
	AMIMIGGIIC	AACCCTATT	GTGAAGAAA	ATGCTTCGGA	2080
	ATTAITICCA	CTTAACGTGA	TAAGTGGAC	AATCTTCGAC	2120
	ተልጥሮልሮሞልሞር	ATGGCTTACA	TGACACAGA	A GACAAAATAA	2160
	DACAGTACGT	GGAAGGCAGI	TCCATTCCT	TTCCAACTCA	2200
25	CTACTACAGC	ATCATCACCA	A GCTGTCTGG	A TTTCACTCAG	2240
	CCTGCCGACA	AGTGTGACG	CCCTCTCTC'	r gtgtcctcct	2280
	TCATCCTGCC	TCACCGGCCT	r GACAAAGAG	G AGAGCTGCAA	2320
	TAGCTCAGAG	GACGAATCA	A AATGGGTAG	A AGAACTCATG	2360
	AAGATGCACA	CAGCTAGGG	r gcgtgacat	I GAACATCTCA	2400
	CCAGCCTGGA	CTTCTTCCG	A AAGACCAGC	C GCAGCTACCC	2440
	AGAAATCCTG	ACACTCAAG	A CATACCTGC	A TACATATGAG	2480
30	AGCGAGATTT	AACTTTCTG	A GCATCTGCA	G TACAGTCTTA	2520
	TCAACTGGTT	GTATATTTT	r atattgttt	T TGTATTTATT	2560
	AATTTGAAAC	CAGGACATT	A AAAATGTTA	G TATTTTAATC	2600
	CTGTACCAAA	TCTGACATA	T TATGCCTGA	A TGACTCCACT	2640
	GTTTTTCTCT	AATGCTTGA	T TTAGGTAGC	C TTGTGTTCTG	2680
	AGTAGAGCTT	GTAATAAAT.	A CTGCAGCTT	G AGAAAAGTG	2720
	GAAGCTTCTA	AATGGTGCT	G CAGATTTGA	T ATTTGCATTG	2760
35	AGGAAATATI	ADDTTTTAA 1	A TGCACAGTT	G CCACATTTAG	2800
33	TCCTGTACTC	3 TATGGAAAC	A CTGATTTG	T AAAGTTGCCT	2840

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	TTAT GCCT AGTT	TATA	AA C	CAAT	CTTA	A AC	ATA	TAAA						2880 2920 2946
	(2)	INF	ORMA	TION	FOR	SEQ	ID	NO:3	6 :					
5		(i)	·	(A) (B) (C)	UENC LE TY ST TC	NGTH PE: RAND	: 7 ami EDNE	88 no a SS:	cid sin	gle				
		(ii	.)	MOL	ECUL	E TY	PE:	pro	tein	ı				
10		(ii	.i)	HYP	OTHE	TICA	L: N	oi						
15		(vi	.)	(A) (B) (C) (D) (E) (F)	OF CE	GANI RAIN DIVI VELO APLOT	SM: I: DUAL OPMEN TYPE: TYPE:	Hum ISC TTAL PE:	LATE	E:	cinom	ıa		
20		(ia	c)	FEA (A) (B)	ATURI NZ LO II O	: AME/H CATI CENTI CHER	ŒY: ION:	ATION	MET	CHOD:			puta	tive
		(xi	L)	SEÇ	QUEN	CE DI	ESCRI	IPTIC	ON: S	SEQ I	ID NO	36:		
25	Cys 1	Asp	Asn	Leu	Cys 5	Lys	Ser	Tyr	Thr	Ser 10	Cys	Cys		
				Asp	Glu					Thr	Ala	Arg		
	Ala 25			Cys						_	Glu 35	Val		
		Asn	Glu	Glu 40	Asn	Ala	Cys	His	Cys 45	Ser		Asp	٠	
30	Cys	Leu 50	Ala	Arg	Gly	Asp	Cys 55	Cys		Asn	Tyr	Gln 60	٠	
	Val		Cys	Lys	Gly 65	Glu		His	Trp	Val	Asp			
	Asp	Cys	Glu 75	Glu		Lys	Ala	Ala 80	Glu		Leu	Gln		
25	Val 85	Asp		Pro	Ser	Ile 90	Asn		Leu	Leu	Arg 95	Gly		

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•			Pro	חחה					TOO			
	-	310	Met	Pro			115					120
		Thr	His		125					T20		
_			Thr 135					140				
5	115		Tyr			150					T22	
	Ser		Tyr	160					TOD			
		170	Gly				1/5					100
10			Gln		185					TOO		
			Glu 195					200				
	20E		Ser			210					213	
			Thr	220					225			
15		230	Phe				235	)				240
		-	His		245					250		
			Arg 255 Gly	5				260	}			
20	255	:	n Val			270	)				2/5	
20			n val s Met	280	)				28:	)		
		20					29	<b>o</b> .		•		300
			r Asi u Gly		305	5				210		
25			211	5				321	U			Leu
	~ ~	_				221	n.				222	Tyr
				34(	0				54	<b>ગ</b>		Ala
		35	.0				35	5				360 ı Val
30					36	5				3/0	,	ı Asp
			37	5				38	0			r Arg
	3,8	5				39	0				39:	5 r Met
			_	40	0				40	5		e Lys
35	GI		10				43	1.5	•			420

	Tyr	Lys	Thr	Lys	Val 425	Pro	Pro	Phe	Glu	Asn 430	Ile	Glu
	Leu	Tyr	Asn 435	Val	Met	Cys	Asp	Leu 440	Leu	Gly	Leu	Lys
	Pro 445	Ala	Pro	Asn	Asn	Gly 450	Thr	His	Phe	Ser	Leu 455	Asn
5				460	Thr				465			
	Pro	Glu 470	Glu	Val	Thr	Arg	Pro 475	Asn	Tyr	Pro	Gly	Ile 480
		_			Ser 485	_		-		490	-	
	Cys	Asp	Asp 495	Lys	Val	Glu	Pro	Lys 500	Asn	Lys	Leu	Asp
10	Glu 505	Leu	Asn	Lys	Arg	Leu 510	His	Thr	Lys	Gly	Ser 515	Thr
	Glu	Glu	Arg	His 520	Leu	Leu	Tyr	Gly	Asp 525	Arg	Pro	Ala
	Val	Leu 530	Tyr	Arg	Thr	Arg	Tyr 535	Asp	Ile	Leu	Tyr	His 540
15		_			Ser 545	_	-			550		
13	•		555		Thr			560			-	
	565				Ser	570					<b>57</b> 5	
	_		_	580	Asp		_		585			
20		590			Leu		595			_	_	600
					Gly 605		_			610		
			615		Ala			620				,
	625				Pro	630					635	
25				640	Phe				645		-	-
		650			Arg		655					660
					Asp 665				_	670		
20			675		Lys Val			680				-
30	685				Leu	690					695	
				700	Pro				705			
		710			Pro		715					720
35	Tien	FIO	nis	wrA	725	nah	nali	GIU	GIU	730	Cys	ASN

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Ser Ser Glu Asp Glu Ser Lys Trp Val Glu Glu Leu 740 735 Met Lys Met His Thr Ala Arg Val Arg Asp Ile Glu 750 His Leu Thr Ser Leu Asp Phe Phe Arg Lys Thr Ser 765 760 Arg Ser Tyr Pro Glu Ile Leu Thr Leu Lys Thr Tyr 775 5 Leu His Thr Tyr Glu Ser Glu Ile INFORMATION FOR SEQ ID NO:37: (2) SEQUENCE CHARACTERISTICS: (i) LENGTH: 2712 10 (A) TYPE: nucleic acid (B) STRANDEDNESS: double (C) TOPOLOGY: Unknown (D) (ii) MOLECULE TYPE: cDNA HYPOTHETICAL: No (iii) 15 ORIGINAL SOURCE: (vi) ORGANISM: Human (A) STRAIN: (B) INDIVIDUAL ISOLATE: (C) DEVELOPMENTAL STAGE: (D) (E) HAPLOTYPE: (F) TISSUE TYPE: 20 CELL TYPE: teratocarcinoma (G) CELL LINE: N-tera 2D1 (H) ORGANELLE: (I) (ix) FEATURE: NAME/KEY: (A) (B) LOCATION: 25 IDENTIFICATION METHOD: (C) OTHER INFORMATION: N-tera 2D1 ATX DNA (D) sequence SEQUENCE DESCRIPTION: SEQ ID NO:37: (xi) 40 TGTGACAACT TGTGTAAGAG CTATACCAGT TGCTGCCATG ACTTTGATGA GCTGTGTTTG AAGACAGCCC GTGCGTGGGA 80 30 GTGTACTAAG GACAGATGTG GGGAAGTCAG AAATGAAGAA 120 160 AATGCCTGTC ACTGCTCAGA GGACTGCTTG GCCAGGGGAG ACTGCTGTAA CAATTACCAA GTGGTTTGCA AAGGAGAGTC 200 -GCATTGGGTT GATGATGACT GTGAGGAAAT AAAGGCCGCA 240 GAATGCCTGC AGGTTTGTTC GCCCTCCATT AATCATCTTC 280 TCCGTGGATG GCTTCCGATG ACATCATACA TGAAGAAAGG 320 CAGCAAAGTC ATGCCTAATA TTGAAAAACT AAGGTCTTGT 360 GGCACACACT CTCCCTACAT GAGGCCGGTG TACCCAACTA 400 35

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•					
	AAACCTTTCC	TAACTTATAC	ACTTTGGCCA	CTGGGCTATA	440
			TTGGCAATTC		480
			TCATCTGCGA		520
	AATTTAATCA	TAGATGGTGG	GGAGGTCAAC	CGCTATGGAT	560
			GTGAAAGCTG		600
	TTGGTCTGTT	GTCATCCCTC	ACGAGCGGAG	ATATTAACCA	640
5			CCAGATCATG		680
,	GGTCTATGCC	TTCTATTCTG	AGCAACCTGA	TTTCTCTGGA	720
			CCCTGAGATG		760
	TGAGGGAAAT	GCACAAAATT	GTGGGGCAAT	TAATGGATGG	800
	ACTGAAACAA	CTAAAACTGC	ATCGGTGTGT	CAACGTCATC	840
	TTTGTCGAGA	CCATGGATGG	AAGATGTCAC	ATGTATAGAA	880
	CTGAGTTCTT	GAGTAATTAC	CTAACTAATG	TGGATGATAT	920
	TACTTTAGTG	CCTGGAACTC	TAGGAAGAAT	TCGATCCAAA	960
10	TTTAGCAACA	ATGCTAAATA	TCACCCCAAA	GCCATTATTG	1000
	CCAATCTCAC	GTGTAAAAA	CCAGATCAGC	ACTTTAAGCC	1040
	TTACTTGAAA	CAGCACCTTC	CCAAACGTTT	GCACTATGCC	1080
	AACAACAGAA	GAATTGAGGA	TATCCATTTA	TTGGTGGAAC	1120
	GCAGATGGCA	TGTTGCAAGG	AAACCTTTGG	ATGTTTATAA	1160
	GAAACCATCA	GGAAATGCTT	TTTCCAGGGA	GACCACGGCA	1200
	TTTGATAACA	AGGTCAACAG	CATGCAGACT	GTTTTTGTAG	1240
15	GTTATGGCCC	AACATTTAAG	TACAAGACTA	AAGTDCCTCC	1280
1.5			AAAATGTTAT		1320
			TAATAATGGG		1360
	GTTTGAATCA	TCTCCTGCGC	ACTAATACCT	TCAGGCCAAC	1400
			GACCCTATTA		1440
			TGACCTGGGC		1480
			AACAAGTTGG		1520
20			GGTCTACAGA		1560
20			TGCAGTGCTT		1600
			ACTGACTTTG		1640
			CACTCTGGAC TTCCAGCGTT		1680 1720
			GATGTCCGTG		1760
			CCTACAAAA		1800
			TCCTCCTTAT		1840
			GCATTCCTTG		1880
25			TCAAACGGGT		1920
			GAAATATGCT		1960
			GGACCAATCT		2000
			CAGAAGACAA		2040
			TCCTGTTCCA		2080
			CTGGATTTCA		2120
			TCTCTGTGTC		2160
30			CGAGGAGAGC		2200
			GTAGAAGAAC		2240
	GCACACAGCT	AGGGTGCGTG	ACATTGAACA	TCTCACCAGC	2280
	CTGGACTTCT	TCCGAAAGAC	CAGCCGCAGC	TACCCAGAAA	2320
			CTGCATACAT		2360
			TGCAGTACAG		2400
			GTTTTTGTAT		2440
35			GTTAGTATTT		2480
J	CCAAATCTGA	CATATTATGC	CTGAATGACT	CCACTGTTTT	2520

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0		-												
•	TCTCT AGCTT CTTCT AATAT GTACT	GTAA 'AAAT Taatu	T AA G GT T TT	ATAC GCTG CCAA	TGCA CAGA TGCA	GCT TTT CAG	TGAG GATA TTGC	TTT TTT CAC	TTAG GCAT ATTT	TGAG	AG GA		2560 2600 2640 2680 2712	
5	(2) INFORMATION FOR SEQ ID NO:38:													
		(i)		(A) (B) (C)		IGTH : PE : RANDE	97 amir EDNES	79 no ac SS:	id sing					
10		(ii)	)	MOLI	ECULI	E TYI	E:	prot	ein					
		(ii:	i)	HYPO	OTHE:	[[CA]	ւ: No	>						
15		(vi	)	(A) (B) (C) (D) (E) (F)	STI INI DE HA TI CE CE	GANIS RAIN DIVII VELO PLOT SSUE LL T	SM: : DUAL PMEN YPE: TYP YPE:	Huma ISO TAL	LATE STAG	E:		-		
20	(ix)			(A) (B) (C) (D)	OT pr	ME/K CATI ENTI HER otei	ON: FICA INFO	RMAT	'ION: ace f	pu rom	tati huma	n liv	totaxin er	
25		(xi	.)	SEÇ	OUENC	CE DE	ESCRI	PTIC	N: S	SEQ I	D NC	):38:		
	1				Ser 5					ΤO				
	Ile	Ser	Leu 15	Phe	Thr	Phe	Ala	Val 20	Gly	Val	Asn	Ile		
20	Cys 25	Leu	Gly	Phe	Thr	Ala 30	His	Arg	Ile	Lys	Arg 35	Ala		
30	Glu	Gly	Trp	Glu 40	Glu		Pro	Pro	Thr 45	Val	Leu	Ser		
		50		Trp	Thr		55		Gly			60		
	Gly	Arg	Cys	Phe	Glu 65	Leu	Gln	Glu	Ala	Gly 70	Pro	Pro		
35	Asp	Cys	Arg 75	Cys	Asp	Asn	Leu	Суs 80	Lys	Ser	Tyr	Thr		

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	Ser 85	Cys	Cys	His	Asp	Phe 90	Asp	Glu	Leu	Cys	Leu 95	Lys
	Thr	Ala	Arg	Ala 100	Trp	Glu	Cys	Thr	Lys 105	Asp	Arg	Cys
	Gly	Glu 110	Val		Asn	Glu	Glu 115	Asn	Ala	Cys	His	Cys 120
5	Ser	Glu	Asp	Cys	Leu 125	Ala		Gly	Asp	Cys 130	Cys	Thr
	Asn	Tyr	Gln 135	Val	Val	Cys	Lys	Gly 140	Glu		His	Trp
	Val 145	Asp	Asp	Asp	Cys	Glu 150	Glu		Lys	Ala	Ala 155	Glu
	Cys	Leu	Gln	Val 160	Cys		Pro	Ser	Ile 165	Asn	His	Leu
10	Leu	Arg 170	Gly		Leu	Pro	Met 175	Thr	Ser	Tyr	Met	Lys 180
	Lys		Ser	Lys	Val 185	Met	Pro	Asn	Ile	Glu 190	Lys	Leu
	Arg	Ser	Cys 195	Gly		His	Ser	Pro 200	Tyr	Met	Arg	Pro
1.5	205	_	Pro			210					Tyr 215	
15	Leu			220					225		Gly	
		230					235				Asp	240
					245					250	Asn	
20			255					260			Thr	
	265					270					Leu 275	
		_		280					285		Leu	
		290					295				Glu	300
25				_	305					310		
			315					320			Pro	
	325					330					335	
	_			340					345		-	Leu
30		350	)				355	;				Met 360
	_				365	•				370	)	Leu
			375	•				380	)			Leu
35	Val 385		o Gly	Thr	Leu	390		i ite	: Arg	ser	туs 395	Phe

o	Ser	Asn	Asn	Ala 400	Lys	Tyr	Asp	Pro	Lys 405	Ala	Ile	Ile
		410	Leu	Thr			415					420
	_	Pro	Tyr		425					430		
<b>.</b>			Ala 435					440				
5	445		Val			450					455	
	Pro		Asp	460					465			
		470	Arg				475					480
10			Met		485					490		
			Lys 495					500				
	505		Glu			510					515	
			Lys	520					525			
15		530	Asn				535					540
			Met		545					550		
			7 Ile 555			-		560				
	565		Thr			570					575	
20			Asp	580					585			
	_	590	Thr				595	,				600
			Ala		605					610		
25			615	;				620	)			Glu
23	625	5				630	)				635	
		_		640	)				645	5		His
		650	0				655	5				Ser 660
30					665	5				670	)	Asn
			675	5				680	0			Pro
	68	5				690	)				695	
				700	0				70!	5		Ala
35	Ph	e Ly 71		g Val	ı Trj	o Asi	71	r Pn 5	e G11	n Arç	y val	720

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```
Val Lys Lys Tyr Ala Ser Glu Arg Asn Gly Val Asn
     Val Ile Ser Gly Pro Ile Phe Asp Tyr Asp Tyr Asp
                                  740
     Gly Leu His Asp Thr Glu Asp Lys Ile Lys Gln Tyr
                          750
     Val Glu Gly Ser Ser Ile Pro Val Pro Thr His Tyr
5
                  760
                                       765
     Tyr Ser Ile Ile Thr Ser Cys Leu Asp Phe Thr Gln
                              775
     Pro Ala Asp Lys Cys Asp Gly Pro Leu Ser Val Ser
                      785
     Ser Phe Ile Leu Pro His Arg Pro Asp Asn Glu Glu
              795
10
     Ser Cys Asn Ser Ser Glu Asp Glu Ser Lys Trp Val
     805
                          810
                                               815
     Glu Glu Leu Met Lys Met His Thr Ala Arg Val Arg
     Asp Ile Glu His Leu Thr Ser Leu Asp Phe Phe Arg
                              835
          830
     Lys Thr Ser Arg Ser Tyr Pro Glu Ile Leu Thr Leu
                      845
15
     Lys Thr Tyr Leu His Thr Tyr Glu Ser Glu Ile Xaa
               855
                                   860
     Leu Ser Glu His Leu Gln Tyr Ser Leu Ile Asn Trp
                          870
     Leu Tyr Ile Phe Ile Leu Phe Leu Tyr Leu Leu Ile
                  880
                                       885
      Xaa Asn Gln Asp Ile Lys Asn Val Ser Ile Leu Ile
20
                               895
      Leu Tyr Gln Ile Xaa His Ile Met Pro Glu Xaa Leu
                      905
     His Cys Phe Ser Leu Met Leu Asp Leu Gly Ser Leu
                                   920
      Val Phe Xaa Val Glu Leu Val Ile Asn Thr Ala Ala
                          930
      Xaa Val Phe Ser Gly Ser Phe Xaa Met Val Leu Gln
25
                  940
                                       945
      Ile Xaa Tyr Leu His Xaa Gly Asn Ile Asn Phe Pro
                               955
      Met His Ser Cys His Ile Xaa Ser Cys Thr Val Trp
                      965
      Lys His Xaa Phe Cys Lys Val
              975
30
```

### (2) INFORMATION FOR SEQ ID NO:39:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 8
  - (B) TYPE: amino acids
  - (C) STRANDEDNESS: single

WO 95/32221 PCT/US95/06613

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(D) TOPOLOGY: linear MOLECULE TYPE: (ii) (A) DESCRIPTION: peptide (iii) HYPOTHETICAL: No (ix) FEATURE: 5 (A) NAME/KEY: ATX-204 (B) LOCATION: IDENTIFICATION METHOD: (C) OTHER INFORMATION: (D) SEQUENCE DESCRIPTION: SEQ ID NO:39: (xi) 10 Met His Thr Ala Arg Val Arg Asp INFORMATION FOR SEQ ID NO:40: (2) SEQUENCE CHARACTERISTICS: (i) (A) LENGTH: 8 15 (B) TYPE: amino acid (C) STRANDEDNESS: single TOPOLOGY: linear (D) (ii) MOLECULE TYPE: peptide HYPOTHETICAL: No (iii) 20 (ix) FEATURE: NAME/KEY: ATX-205 (A) (B) LOCATION: (C) IDENTIFICATION METHOD: (D) OTHER INFORMATION: 25 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:40: Phe Ser Asn Asn Ala Lys Tyr Asp 5 (2) INFORMATION FOR SEQ ID NO:41: (i) 30 SEQUENCE CHARACTERISTICS: (A) LENGTH: 7 TYPE: amino acids (B) STRANDEDNESS: single (C) TOPOLOGY: linear (D) MOLECULE TYPE: (ii) (A) DESCRIPTION: Peptide 35

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D			
		(iii)	HYPOTHETICAL: No
5		(ix)	FEATURE: (A) NAME/KEY: ATX-209 (B) LOCATION: (C) IDENTIFICATION METHOD: (D) OTHER INFORMATION:
		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:41:
	Val 1	Met Pro	Asn Ile Glu Lys 5
10	(2)	INFORMA	ATION FOR SEQ ID NO:42:
		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 8  (B) TYPE: amino acids  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear
15		(ii)	MOLECULE TYPE: (A) DESCRIPTION: Peptide
		(iii)	HYPOTHETICAL: No
20		(ix)	FEATURE: (A) NAME/KEY: ATX-210 (B) LOCATION: (C) IDENTIFICATION METHOD: (D) OTHER INFORMATION:
		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:42:
25	Thr	Ala Arg	Gly Trp Glu Cys Thr
	(2)	INFORM	MATION FOR SEQ ID NO:43:
30		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 11  (B) TYPE: amino acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear
		(ii)	MOLECULE TYPE: (A) DESCRIPTION: Peptide
35		(iii)	HYPOTHETICAL: No

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	(ix)	FEATURE: (A) NAME/KEY: ATX-212 (B) LOCATION: (C) IDENTIFICATION METHOD: (D) OTHER INFORMATION:
5	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:43:
	Xaa Asp Ser I	Pro Trp Thr Xaa Ile Ser Gly Ser 5 10
10	(2) INFORMA	TION FOR SEQ ID NO:44:
10	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 11  (B) TYPE: amino acids  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear
15	(ii)	MOLECULE TYPE: (A) DESCRIPTION: Peptide
	(iii)	HYPOTHETICAL: No
20	(ix)	FEATURE: (A) NAME/KEY: ATX-214 (B) LOCATION: (C) IDENTIFICATION METHOD: (D) OTHER INFORMATION:
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:44:
25	Leu Arg Ser	Cys Gly Thr His Ser Pro Tyr Met 5 10
	(2) INFORMA	TION FOR SEQ ID NO:45:
30	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 8  (B) TYPE: amino acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear
	(ii)	MOLECULE TYPE: (A) DESCRIPTION: Peptide
	(iii)	HYPOTHETICAL: No
35	(ix)	FEATURE:

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0	•	·
		(A) NAME/KEY: ATX-215/34A (B) LOCATION: (C) IDENTIFICATION METHOD: (D) OTHER INFORMATION:
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:45:
5	Thr Tyr Leu	His Thr Tyr Glu Ser 5
	(2) INFORMA	TION FOR SEQ ID NO:46:
10	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 13  (B) TYPE: amino acids  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear
	(ii)	MOLECULE TYPE: (A) DESCRIPTION: Peptide
15	(iii)	HYPOTHETICAL: No
	(ix)	FEATURE: (A) NAME/KEY: (B) LOCATION: (C) IDENTIFICATION METHOD: (D) OTHER INFORMATION:
20	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:46:
	Ala Ile Ile	Ala Asn Leu Thr Cys Lys Lys Pro Asp Gln 5
25	(2) INFORM	ATION FOR SEQ ID NO:47:
	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 8  (B) TYPE: amino acids  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear
30	(ii)	MOLECULE TYPE: (A) DESCRIPTION: Peptide
	· (iii)	HYPOTHETICAL: No
	(ix)	FEATURE: (A) NAME/KEY: ATX-216 (B) LOCATION:
35		(C) IDENTIFICATION METHOD:

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OTHER INFORMATION: (D) SEQUENCE DESCRIPTION: SEQ ID NO:47: (xi) Ile Val Gly Gln Leu Met Asp Gly 5 INFORMATION FOR SEQ ID NO:48: (2) SEQUENCE CHARACTERISTICS: (i) (A) LENGTH: 9 TYPE: amino acids (B) STRANDEDNESS: single (C) TOPOLOGY: linear 10 (D) MOLECULE TYPE: (ii)(A) DESCRIPTION: Peptide HYPOTHETICAL: No (iii) FEATURE: (ix) 15 NAME/KEY: ATX-218/44 (A) LOCATION: (B) IDENTIFICATION METHOD: (C) OTHER INFORMATION: (D) SEQUENCE DESCRIPTION: SEQ ID NO:48: (xi) Thr Ser Arg Ser Tyr Pro Glu Ile Leu 20 5 INFORMATION FOR SEQ ID NO:49: SEQUENCE CHARACTERISTICS: (i) 25 LENGTH: 9 (A) TYPE: amino acids (B) STRANDEDNESS: single (C) TOPOLOGY: linear (D) MOLECULE TYPE: (ii) DESCRIPTION: Peptide (A) 30 HYPOTHETICAL: No (iii) FEATURE: (ix) NAME/KEY: ATX-223B/24 (A) LOCATION: (B)

(C)

(D)

IDENTIFICATION METHOD:

OTHER INFORMATION:

35

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(xi)
                   SEQUENCE DESCRIPTION: SEQ ID NO:49:
     Gln Ala Glu Val Ser Ser Val Pro Asp
5
      (2)
          INFORMATION FOR SEO ID NO:50:
           (i)
                   SEQUENCE CHARACTERISTICS:
                   (A)
                        LENGTH: 14
                   (B)
                        TYPE: amino acids
                        STRANDEDNESS: single
                   (C)
                        TOPOLOGY: linear
                   (D)
10
           (ii)
                   MOLECULE TYPE:
                   (A)
                       DESCRIPTION: Peptide
           (iii)
                  HYPOTHETICAL: No
           (ix)
                   FEATURE:
                   (A)
                        NAME/KEY: ATX-224
                   (B)
                        LOCATION:
15
                   (C)
                        IDENTIFICATION METHOD:
                   (D)
                        OTHER INFORMATION:
           (xi)
                   SEQUENCE DESCRIPTION: SEQ ID NO:50:
     Arg Cys Phe Glu Leu Gln Glu Ala Gly Pro Pro Asp Asp Cys
20
      (2)
          INFORMATION FOR SEO ID NO:51:
           (i)
                   SEQUENCE CHARACTERISTICS:
                   (A)
                        LENGTH:
                                12
                   (B)
                        TYPE: amino acid
                   (C)
                        STRANDEDNESS: single
25
                   (D)
                        TOPOLOGY: linear
           (ii)
                   MOLECULE TYPE:
                   (A) DESCRIPTION: Peptide
           (iii)
                   HYPOTHETICAL: No
30
           (ix)
                   FEATURE:
                   (A)
                        NAME/KEY: ATX-229
                        LOCATION:
                   (B)
                   (C)
                        IDENTIFICATION METHOD:
```

OTHER INFORMATION:

SEQUENCE DESCRIPTION: SEQ ID NO:51:

(D)

(xi)

BNSDOCID: <WO\_\_9532221A2\_I\_>

35

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Ser Tyr Thr Ser Cys Cys His Asp Phe Asp Glu Leu
          INFORMATION FOR SEQ ID NO:52:
     (2)
                  SEQUENCE CHARACTERISTICS:
          (i)
                       LENGTH: 16
                   (A)
5
                       TYPE: amino acid
                   (B)
                        STRANDEDNESS: single
                   (C)
                        TOPOLOGY: linear
                   (D)
                  MOLECULE TYPE:
          (ii)
                   (A) DESCRIPTION: Peptide
                   HYPOTHETICAL: No
10
          (iii)
                   FEATURE:
           (ix)
                       NAME/KEY: ATX-224/53
                   (A)
                        LOCATION:
                   (B)
                        IDENTIFICATION METHOD:
                   (C)
                        OTHER INFORMATION:
                   (D)
15
                   SEQUENCE DESCRIPTION: SEQ ID NO:52:
           (xi)
     Gln Met Ser Tyr Gly Phe Leu Phe Pro Pro Tyr Leu
                        5
      Ser Ser Ser Pro
               15
20
           INFORMATION FOR SEQ ID NO:53:
      (2)
                   SEQUENCE CHARACTERISTICS:
           (i)
                       LENGTH: 117
                    (A)
                        TYPE: nucleic acid
                    (B)
                        STRANDEDNESS: single
                    (C)
                        TOPOLOGY: Unknown
                    (D)
25
                   MOLECULE TYPE:
            (ii)
                    (A) DESCRIPTION:
                                        cDNA
                   HYPOTHETICAL: No
            (iii)
                    ANTI-SENSE:
            (iv)
 30
                    ORIGINAL SOURCE:
            (vi)
                    (A) ORGANISM: Human
                    (B) STRAIN:
                         INDIVIDUAL ISOLATE:
                    (C)
                     (D) DEVELOPMENTAL STAGE:
                         HAPLOTYPE:
                     (E)
                         TISSUE TYPE: Liver
                     (F)
 35
                         CELL TYPE:
                     (G)
```

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0		
		(H) CELL LINE: (I) ORGANELLE:
5	(ix)	FEATURE: (A) NAME/KEY: (B) LOCATION: (C) IDENTIFICATION METHOD: (D) OTHER INFORMATION: 5' end of human liver ATX gene
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:53:
10	CCCTGTTCAC	GGAGCTCGTT CCAGTCGTGT CAAGATATAT 40 TTTTGCCGTT GGAGTCAATA TCTGCTTAGG 80 CATCGAATTA AGAGAGCAGA AGGATGG 117
	(2) INFOR	MATION FOR SEQ ID NO:54:
15	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 39 (B) TYPE: amino acids (C) STRANDEDNESS: single (D) TOPOLOGY: Unknown
	(ii)	MOLECULE TYPE: (A) DESCRIPTION: Peptide
	(iii)	HYPOTHETICAL: No
20	(v)	FRAGMENT TYPE: N-terminal fragment
25	(vi)	ORIGINAL SOURCE: (A) ORGANISM: Human (B) STRAIN: (C) INDIVIDUAL ISOLATE: (D) DEVELOPMENTAL STAGE: (E) HAPLOTYPE: (F) TISSUE TYPE: Liver (G) CELL TYPE: (H) CELL LINE: (I) ORGANELLE:
30	(ix)	FEATURE:  (A) NAME/KEY: (B) LOCATION: (C) IDENTIFICATION METHOD: (D) OTHER INFORMATION: N-terminal region including transmembrane domain of liver ATX protein
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:54:
35		

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•										
•		la Arg	Arg Ser		e Gln	Ser	Cys 10	Gln	Asp	
	lle Se	er Leu 15	Phe Thr		a Val 20	Gly		Asn	Ile	
			Phe Thr	Ala Hi		Ile	Lys	Arg 35	Ala	
	25 Glu G	ly Trp		30				23		
5										
	(2)	INFORM	ATION FO	R SEQ I	D NO:	55:				
		(i)		ICE CHAR		ISTI	CS:			
			(B) 7		21 mino					
10				TRANDED TOPOLOGY				-		
		(ii)		JLE TYPE						
		(iii)	нүроті	HETICAL:	No					
15		(iv)	ANTI-	SENSE:	Yes					
		(ix)	FEATU							
				NAME/KEY LOCATION					-	
	•			IDENTIF		N ME	THOD	):		
20				OTHER IN 4C11	IFORM <i>I</i>	MOITA	I: Pr	rimer	from 5	' end of
		(xi)	SEQUE	NCE DESC	RIPT	ON:	SEQ	ID N	10:55:	
	GCTCA	GATAA	GGAGGAA	AGA G						2
25	(2)	INFORM	ATION F	OR SEQ :	D NO	:56:				
23		(i)	SEQUE	NCE CHAI	RACTE	RIST	cs:			
			(A)	LENGTH: TYPE: 8	21					
			(C)	STRANDE	ONESS	: si	ingle	2		
				TOPOLOG		inear	5			
30		(ii)	MOLEC	ULE TYP	E: cD	AV				
		( <b>iii</b> )	HYPOT	HETICAL	: No					
		(iv)	ANTI-	SENSE:	Yes					
		(ix)	FEATU		σ.					
35				NAME/KE LOCATIO						

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•	•				
			(C) (D)	IDENTIFICATION METHOD: OTHER INFORMATION: Nested primers from 4C11	ı
		(xi)	SEQU	ENCE DESCRIPTION: SEQ ID NO:56:	
5	GAAT	CCGTAG G	ACATC	TGCT T	21
	(2)	INFORMA	MOITA	FOR SEQ ID NO:57:	
10		(i)	(A) (B) (C)	WENCE CHARACTERISTICS: LENGTH: 21 TYPE: amino acid STRANDEDNESS: single TOPOLOGY: linear	
		(ii)	MOLE	ECULE TYPE: cDNA	
		(iii)	НҮРС	OTHETICAL: No	
15		(iv)	ANTI	I-SENSE: Yes	
		(ix)	(A) (B) (C)	TURE: NAME/KEY: LOCATION: IDENTIFICATION METHOD: OTHER INFORMATION: Nested primers fro 4C11	m
20		(xi)	SEQ	UENCE DESCRIPTION: SEQ ID NO:57:	
	TGT	AGGCCAA	ACAGT'	TCTGA C	21
	(2)	INFORM	ATION	FOR SEQ ID NO:58:	
25		(i)	SEQ (A) (B) (C) (D)	STRANDEDNESS: single	
		(ii)	MOL	ECULE TYPE: cDNA	
30		(iii)	НУР	OTHETICAL: No	
		(iv)	ANT	'I-SENSE: No	
35		(ix)	FEA (A) (B)	LOCATION:	

PCT/US95/06613

WO 95/32221

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			(D) OTHER INFORMATION: Nested sense primer deduced from ATX-101, wherein N is inosine	
		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:58:	
5 A	AYTC	NATGC AR	PACNGINIT YGING	25
(	(2)	INFORMAT	TION FOR SEQ ID NO:59:	
10		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 26  (B) TYPE: amino acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
		(ii)	MOLECULE TYPE: cDNA	
		(iii)	HYPOTHETICAL: No	
15		(iv)	ANTI-SENSE: No	
		(ix)	FEATURE: (A) NAME/KEY: (B) LOCATION: (C) IDENTIFICATION METHOD: (D) OTHER INFORMATION: Nested primer of AT -101, wherein N is inosine	<b>'X</b>
20		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:59:	
•	TTYG	INGGNT A	YGGNCCNAC NTTYAA	26
	(2)	INFORMĄ	TION FOR SEQ ID NO:60:	
25		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 26  (B) TYPE: amino acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	-
		(ii)	MOLECULE TYPE: cDNA	
30		(iii)	HYPOTHETICAL: No	
		(iv)	ANTI-SENSE: No	
35		(ix)	FEATURE: (A) NAME/KEY: (B) LOCATION: (C) IDENTIFICATION METHOD:	

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•				
,			(D) OTHER INFORMATION: Nested primer deduction from ATX-103, wherein N is inosine	ed:
		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:60:	
5	AAYT.	AYCTNA (	CNAAYGTNGA YGAYAT	26
	(2)	INFORM	ATION FOR SEQ ID NO:61:	
10		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 26  (B) TYPE: amino acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
		(ii)	MOLECULE TYPE: cDNA	
		(iii)	HYPOTHETICAL: No	
		(iv)	ANTI-SENSE: No	
15	·	(ix)	FEATURE: (A) NAME/KEY: (B) LOCATION: (C) IDENTIFICATION METHOD: (D) OTHER INFORMATION: Nested primer dedufrom ATX-103, wherein N is inosine	ced
20		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:61:	
	GAYO	SAYATNA	CNCTNGTNCC NGGNAC	26
	(2)	INFORM	MATION FOR SEQ ID NO:62:	
25		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 29  (B) TYPE: amino acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
		(ii)	MOLECULE TYPE: cDNA	
30		(iii)	HYPOTHETICAL: No	
		(iv)	ANTI-SENSE: No	
35		(ix)	FEATURE: (A) NAME/KEY: (B) LOCATION: (C) IDENTIFICATION METHOD:	

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0			(D) OTHER INFORMATION: Nested primer ded from ATX-103, wherein N is inosine	uced
		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:62:	
	TGYT'	TYGARY T	INCARGARGC NGGNCCNCC	29
5	(2)	INFORMA	ATION FOR SEQ ID NO:63:	
10		(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 18 (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
10		(ii)	MOLECULE TYPE: cDNA	
		(iii)	HYPOTHETICAL: No	
		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:63:	
15	GCTG	TCTTCA A	AACACAGC	18
	(2)	INFORM	ATION FOR SEQ ID NO:64:	
20		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 21  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
		(ii)	MOLECULE TYPE: cDNA	
		(iii)	HYPOTHETICAL: No	
25		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:64:	
	CTG	GTGGCTG	TAATCCATAG C	21
	(2)	INFORM	MATION FOR SEQ ID NO:65:	
30		(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 21  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
		(ii)	MOLECULE TYPE: cDNA	
25	•	(iii)	HYPOTHETICAL: No	

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_								
		(iv)	ANTI-	SENSE:	No			
5		(ix)	(B) (C) (D)	NAME/KEY LOCATION IDENTIFE	1:  CATION    IFORMATI	ON: Primer	for 5'	end of
		(xi)	SEQUE	NCE DESC	CRIPTION	: SEQ ID N	10:65:	
	CGTGA	AGGCA .	AAGAGAA	CAC G				21
10	(2)	INFORM	ATION F	FOR SEQ	ID NO:66	:		
		(i)	(A) (B)	TYPE: 1	3104 nucleic ONESS:	acid double		
15		(ii)	MOLE	CULE TYP	E: cDNA			
		(iii)	HYPO	THETICAL	: No			
20		(ix)	FEAT( (A) (B) (C) (D)	NAME/KE LOCATION IDENTIF	N:		K cDNA	
		(xi)	SEQU	ENCE DES	CRIPTION	: SEQ ID	NO:66:	
25	CTTTC CAGTC GAGTC GAGAG TCAGA	CAATA GTGTC AATAT CAGAA CTCCC	ATCCTCO AGATAA' CTGCTTA GGATGGO CCTGGAO	GACA TGG IATC CCT AGGA TTC GAGG AAG CCAA CAT	CAAGGAG GTTCACT ACTGCAC GTCCTCC CTCCGGA	CTGCAAAAG GAGCTCGTT TTTGCCGTT ATCGAATTA TACAGTGCT TCTTGCAAG	C G A A G	40 80 120 160 200 240
30	TCGCT CATGA GGGAG AGAAA GGAGA AGTCG	GTGAC CTTTG TGTAC ATGCC CTGCT CATTG	AACTTG ATGAGC TAAGGA TGTCAC GTACCA GGTTGA	IGTA AGA IGTG TTT CAGA TGT TGCT CAG ATTA CCA TGAT GAC	GCTATAC GAAGACA GGAGAAG AGGACTG AGTGGTT TGTGAGG	CTCCTGATT CAGTTGCTG GCCCGTGCG TCAGAAATG CTTGGCCAG TGCAAAGGA AAATAAAGG TCCATTAAT	C T A G G C	280 320 360 400 440 480 520 560
35	ATCTT AAGGC TTGTG	CTCCG CAGCAA GCACA	TGGATG AGTCAT CACTCG	GCTT CCG GCCT AAT CCCC ACA	TGCATCA ATTGAAA TGAGGCC	TACATGAAG AACTAAGGT GGTGTACCC GCCACTGGG	A C A	600 640 680 720

	•		•		
	TATATCCAGA	ATCACATGGA	ATTGTTGGCA	ATTCAATGTA	760 800 840 880 920 960 1000 1040
	TGATCCTGTA	TTTGATGCCA	CTTTTCATCT	GCGAGGGCGA	800
	GAGAAATTTA	ATCATAGATG	GTGGGGAGGT	CAACCGCTAT	840
	<b>GGATTACAGC</b>	CACCAAGCAA	AGGGGTGAAA	GCTGGAACAT	880
	TCTTTTGGTC	TGTTGTCATC	CCTCACGAGC	GGAGATATTA	920
	ACCATATTGC	AGTGGCTCAC	CCTGCCAGAT	CATGAGAGGC	960
5	TTCGGTCTAT	GCCTTCTATT	CTGAGCAACC	TGATTTCTCT	1000
,	GGACACAAAT	ATGCCTTTCG	GCCCTGAGAT	GACAAATCCT	1040
					7770
	CTTTGTCGGA	GACCATGGAA	TGGAAGATGT	CACATGTGAT	1160
	AGAACTGAGT	TCTTGAGTAA	TTACCTAACT	AATGTGGATG	1120 1160 1200 1240 1280 1320 1360 1400
	ATATTACTTT	AGTGCCTGGA	ACTCTAGGAA	TTCGATCCAA	1240
	ATTTAGCAAC	AATGCTAAAT	ATGACCCCAA	AGCCATTATT	1280
10	GCCAATCTCA	CGTGTAAAAA	ACCAGATCAG	CACTTTAAGC	1320
	CTTACTTGAA	ACAGCACCTT	CCCAAACGTT	TGCACTATGC	1360
	CAACAACAGA	AGAATTGAGG	ATATCCATTT	ATTGGTGGAA	1400
	CGCAGATGGC	ATGTTGCAAG	GAAACCTTTG	GATGTTTATA	1440
	AGAAACCATC	AGGAAAATGC	TTTTTCCAGG	GAGACCACGG	1480
	ATTTGATAAC	AAGGTCAACA	GCATGCAGAC	TGTTTTTGTA	1520
	GGTTATGGCC	CAACATTTAA	GTACAAGACT	AAAGTGCCTC	1560
	CATTTGAAAA	CATTGAACTT	TACAATGTTA	TGTGTGATCT	1600
15	CCTGGGATTG	AAGCCAGCTC	CTAATAATGG	GACCCATGGA	1640
	AGTTTGAATC	ATCTCCTGCG	CACTAATACC	TTCAGGCCAA	1400 1440 1480 1520 1560 1600 1640 1680
	CCATGCCAGA	GGAAGTTACC	AGACCCAATT	ATCCAGGGAT	1720
	TATGTACCTT	CAGTCTGATT	TTGACCTGGG	CTGCACTTGT	1720 1760 1800 1840 1880 1920
	GATGATAAGG	TAGAGCCAAA	GAACAAGTTG	GATGAACTCA	1800
	ACAAACGGCT	TCATACAAAA	GGGTCTACAG	AAGAGAGACA	1840
	CCTCCTCTAT	GGGCGACCTG	CAGTGCTTTA	TCGGACTAGA	1880
20	TATGATGTCT	TATATCACAC	TGACTTTGAA	AGTGGTTATA	1920
	GTGAAATATT	CCTAATGCCA	CTCTGGACAT	CATATACTGT	1960
	TTCCAAACAG	GCTGAGGTTT	CCAGCGTTCC	TGACCATCTG	2000
	ACCAGTTGCG	TCCGGCCTGA	TGTCCGTGTT	TCTCCGAGTT	2040
	TCAGTCAGAA	CTGTTTGGCC	TACAAAAATG	ATAAGCAGAT	2080
	GTCCTACGGA	TTCCTCTTTC	CTCCTTATCT	GAGCTCTTCA	1960 2000 2040 2080 2120
	CCAGAGGGTA	DIADIALAA	AIICCIIGIA	VCCWINIGO	2100
25	TTCCAATGTA	TCCTGCTTTC	AAACGGGTCT	GGAATTATTT	2200
25	CCAAAGGGTA	TTGGTGAAGA	AATATGCTTC	GGAAAGAAAT	2240
	GGAGTTAACG	TGATAAGTGG	ACCAATCTTC	GACTATGACT TAAAACAGTA	2280
	ATGATGGCTT	ACATGACACA	GAAGACAAAA	TAAAACAGTA	2320
	CGTGGAAGGC	AGTTCCATTC	CTGTTCCAAC	TCACTACTAC	2360
	AGCATCATCA	CCAGCTGTCT	GGATTTCACT	CAGCCTGCCG	2400
	ACAAGTGTGA	CGGCCCTCTC	TCTGTGTCCT	CCTTCATCCT	2440
	CCGTCACCGG	CCTGACAACG	AGGAGAGCTG	CAATAGCTCA	2480
30				ATGAAGATGC	2520
-	ACACGGCTAG	GGTGCGTGAC	ATTGAACATC	TCACCAGCCT	2560
	GGACTTCTTC	CGAAAGACCA	GCCGCAGCTA	CCCAGAAATC	2600
				GAGAGCGAGA	2640
				TTATCAACTG	2680
				ATTAATTTGA	2720
	AACCAGGACA	TTAAAAATGT	TAGTATTTTA	ATCCTGTACC	2760
25				ACTGTTTTTC	2800
35				CTGAGTAGAG	2840

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0													
5	CTTG' TCTA TATT: ACTG' GCTG TAAA TTCT	AATG( AATT' TATG( TTAA( CCAA'	GT G TT C GA A CT G TC T	CTGC CAAT ACAC TTAA TAAA	AGAT' GCAC' TGAT' CTAT CATA	T TG. A GT T TT G AC. A TA	ATAT' TGCC GTAA AGAT AATC	ITGC ACAT AGTT ATAT	TTAC GCCT	GAGGZ GTCC' I'TTA' AGCC'	AAA PGT PTT PTA		2880 2920 2960 3000 3040 3080 3104
	(2) INFORMATION FOR SEQ ID NO:67:											,	
10		(i)		(A) (B) (C)	LE TY ST	NGTH PE: RAND	: 8 ami EDNE	TERI 61 no a SS: Unk	cid sin	gle			
		(ii	)	MOL	ECUL	E TY	PE:	pro	tein				
		(ii	i)	HYP	OTHE	TICA	L: N	Ю					
15		(ix	:)	(A) (B) (C)	IC	ME/K CATI ENTI	ON:	N-t TION RMAT	MET	HOD:		protein	·
		(sei		SEÇ	NIENIC	זר שי	rscr i	ייים <i>ו</i>	N · S	SEO I	D <b>N</b> C	):67:	
20	Met			Arg						Cys			
	1 Ile	Ser		Phe	5 Thr	Phe	Ala	Val 20	Gly	10 Val	Asn	Ile	
	Cys 25	Leu	15 Gly	Phe	Thr	Ala 30	His		Ile	Lys	Arg 35	Ala	
25	Glu	_	_	Glu 40					45				
	Asp	Ser 50	Pro	Trp	Thr	Asn	Ile 55	Ser	Gly	Ser	Cys	Lys 60	
-	Gly	Arg	Cys	Phe	Glu 65	Leu	Gln	Glu	Ala	Gly 70	Pro	Pro	
	Asp	Cys	Arg 75	Cys		Asn	Leu	Cys 80	Lys	Ser	Tyr	Thr	
30	Ser 85	Cys	Cys	His	Asp	Phe 90	Asp	Glu	Leu	Cys	Leu 95	Lys	
	Thr	Ala	Arg	Ala 100	Trp	Glu	Cys	Thr	Lys 105	Asp	Arg	Cys	
	Gly	Glu 110	Val	Arg	Asn	Glu	Glu 115			Cys	His	Cys 120	
35	Ser		Asp	Cys	Leu 125				Asp	Cys 130	Cys	Thr	

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•	Asn	Tyr	Gln 135	Val	Val	Cys	Lys	Gly 140	Glu	Ser	His	Trp
	Val 145	Asp		Asp	Cys	Glu 150	Glu		Lys	Ala 155	Ala	Glu
		Pro	Ala	Gly 160	Phe	Val	Arg	Pro	Pro 165	Leu	Ile	Ile
5	Phe	Ser 170	Val	Asp	Gly	Phe	Arg 175	Ala	Ser	Tyr	Met	Lys 180
J	Lys	Gly	Ser	Lys	Val 185	Met	Pro	Asn	Ile	Glu 190	Lys	Leu
	Arg	Ser	Cys 195	Gly	Thr	His	Ser	Pro 200	His	Met	Arg	Pro
	Val 205	Tyr	Pro	Thr	Lys	Thr 210	Phe	Pro	Asn	Leu	Tyr 215	Thr
10		Ala		220		_			225			
	Val	Gly 230	Asn	Ser	Met	Tyr	Asp 235	Pro	Val	Phe	Asp	Ala 240
		Phe			245					250		
	_	Trp	255	_	_			260	_			
15	265	Lys				270					275	
-		Cys		280					285			
		Leu 290					295		_			300
20		Arg			305					310		
20		Leu	315					320	_			
	325	Asn -				330		_			335	_
		Leu		340	_		_		345	_		
25	_	Cys 350					355		_	_		360
		Glu	_		365	_	_	_		370		
		Asn	375					380				
	385	Pro				390				_	395	
30		Asn		400	_	_		_	405			
		Leu 410			_	_	415					420
		Tyr			425				-	430		
	_	Ala	435		_			440	-			
35	1445	Val	GIU	Arg	Arg	1rp 450	HIS	vaı	АТА	arg	Lys 455	Pro

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```
Leu Asp Val Tyr Lys Lys Pro Ser Gly Lys Cys Phe
    Phe Gln Gly Asp His Gly Phe Asp Asn Lys Val Asn
                             475
    Ser Met Gln Thr Val Phe Val Gly Tyr Gly Pro Thr
                                         490
                     485
    Phe Lys Tyr Lys Thr Lys Val Pro Pro Phe Glu Asn
5
                                 500
             495
     Ile Glu Leu Tyr Asn Val Met Cys Asp Leu Leu Gly
                         510
     Leu Lys Pro Ala Pro Asn Asn Gly Thr His Gly Ser
                                      525
                 520
     Leu Asn His Leu Leu Arg Thr Asn Thr Phe Arg Pro
                              535
         530
     Thr Met Pro Glu Glu Val Thr Arg Pro Asn Tyr Pro
10
                     545
     Gly Ile Met Tyr Leu Gln Ser Asp Phe Asp Leu Gly
             555
     Cys Thr Cys Asp Asp Lys Val Glu Pro Lys Asn Lys
                         570
     Leu Asp Glu Leu Asn Lys Arg Leu His Thr Lys Gly
                 580
     Ser Thr Glu Glu Arg His Leu Leu Tyr Gly Arg Pro
15
                              595
     Ala Val Leu Tyr Arg Thr Arg Tyr Asp Val Leu Tyr
                      605
     His Thr Asp Phe Glu Ser Gly Tyr Ser Glu Ile Phe
                                  620
              615
     Leu Met Pro Leu Trp Thr Ser Tyr Thr Val Ser Lys
                          630
20
      Gln Ala Glu Val Ser Ser Val Pro Asp His Leu Thr
                                      645
                  640
      Ser Cys Val Arg Pro Asp Val Arg Val Ser Pro Ser
                              655
      Phe Ser Gln Asn Cys Leu Ala Tyr Lys Asn Asp Lys
                      665
      Gln Met Ser Tyr Gly Phe Leu Phe Pro Pro Tyr Leu
25
                                   680
      Ser Ser Ser Pro Glu Ala Lys Tyr Asp Ala Phe Leu
                           690
      Val Thr Asn Met Val Pro Met Tyr Pro Ala Phe Lys
                                       705
                  700
      Arg Val Trp Asn Tyr Phe Gln Arg Val Leu Val Lys
                               715
      Lys Tyr Ala Ser Glu Arg Asn Gly Val Asn Val Ile
 30
                                           730
                       725
      Ser Gly Pro Ile Phe Asp Tyr Asp Tyr Asp Gly Leu
                                   740
               735
      His Asp Thr Glu Asp Lys Ile Lys Gln Tyr Val Glu
                           750
      Gly Ser Ser Ile Pro Val Pro Thr His Tyr Tyr Ser
                   760
 35
```

GIACCARTIA CCRACTOR 520  GGTTGATGAT GACTGTGAGG AAATAAAGGC CGCAGAATGC 520  CCTGCAGGGT TTGTTCGCCC TCCATTAATC ATCTTCTCCG 560  TGGATGGCTT CCGTGCATCA TACATGAAGA AAGGCAGCAA 600														
Asp Lys Cys Asp Gly Pro Leu Ser Val Ser Ser Phe 785   785	• .	Ile		Thr	Ser	Cys	Leu	Asp 775	Phe	Thr	Gln	Pro	Ala 780	
The Leu Arg His Arg Pro Asp Asn Glu Glu Ser Cys 800		Asp	Lys	Cys	Asp	Gly	Pro		Ser	Val	Ser 790	Ser	Phe	
Asn   Ser   Ser   Glu   Asp   Glu   Ser   Lys   Trp   Val   Glu   Glu   815		Ile	Leu		His	Arg	Pro	Asp	Asn 800	Glu	Glu	Ser	Cys	
Leu Met Lys Met His Thr Ala Arg Val Arg Asp 11e 820 820 825 840 825 840 830 836 835 840 835 840 835 840 835 840 850 845 840 850 845 840 850 860 855 860 860 860 860 860 860 860 860 860 860			Ser	Ser	Glu	Asp	Glu	Ser		Trp	Val	Glu 815	Glu	
Glu His Leu Thr Ser Leu Asp Phe Phe Arg Lys Thr 830   Ser Arg Ser Tyr Pro Glu Ile Leu Thr Leu Lys Thr 845   845   850   851	5	Leu	Met	Lys		His		Ala	Arg	Val	Arg	Asp	Ile	
Ser Arg Ser Tyr Pro Glu Ile Leu Thr Leu Lys Thr		Glu		Leu		Ser	Leu	Asp	Phe		Arg	Lys	Thr 840	-
Tyr Leu His Thr Tyr Glu Ser Glu Ile		Ser	Arg	Ser	Tyr		Glu		Leu	Thr	Leu 850	Lys		
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 3251 (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: Unknown  (ii) MOLECULE TYPE: cDNA  (iii) HYPOTHETICAL: No  (ix) FEATURE: (A) NAME/KEY: A2058 ATX cDNA (B) LOCATION: (C) IDENTIFICATION METHOD: (D) OTHER INFORMATION:  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:68:  25 CGTGAAGGCA AAGAGAACAC GCTGCAAAAG GCTTCCAAGA 40 ATCCTCGACAA TGGCAAGGAG GAGCTCGTTC CAGTCGTGTC 80 AGATAATATC CCTGTTCACT TTTGCCGTTG GAGTCAGTAT 120 CTGCTTAGGA TTCACTGCAC ATCGAATTAA GAGCAGAA 160 GGATGGAGGA AAGGTCCTCC TACAGTGTAT TCAGACTCCC 200 CCTGGACCAA CATCTCCGGA TCTTGCAGG GCAGGTGCTT 240 TGAACTTCAA GAGGCTGGAC CTCCTGATTG TCGCTGTGAC 280 AACTTGTGTA AGAGCTCATC CAGTTGCTC CATGACTTTG 320 ATGAGCTGTG TTTGAAGACA GCCCGTGGCT GGGAGTGTAC 360 TGTCACTCT CAGAGACTG CTCAGAATGA AGAAAATGCC 400 TGTCACTACT CAGAGACTG CTCAGAATGA AGAAAATGCC 400 TGTCACTACT CAGAGACTG CTCAGAATGA AGAAAATGCC 400 TGTCACTACT CAGAGACTG CTTGGCCAGG GGAGACTGCT 440 GGTTGATGAT GACCAGTGGTT TGCAAAGGAG AGTCGCATTG 480 GGTTGATGAT GACTGTGAGG AAATAAAAGGC CGCAGAATGC 520 CCTGCAGGGCT TCGTTGCCCC TCCATTAATC ATCTTCTCCCG 560 TGGATGGGCTT CCGTGCTTCA TACATGAAAA AAAGAAATGC	10	Tyr	Leu		Thr		Glu	Ser	Glu 860	Ile				•
(A) LENGTH: 3251 (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: Unknown  (ii) MOLECULE TYPE: cDNA  (iii) HYPOTHETICAL: No  (ix) FEATURE: (A) NAME/KEY: A2058 ATX cDNA (B) LOCATION: (C) IDENTIFICATION METHOD: (D) OTHER INFORMATION:  (Xi) SEQUENCE DESCRIPTION: SEQ ID NO:68:  25 CGTGAAGGCA AAGAGAACAC GCTGCAAAAG GCTTCCAAGA 40 ATCCTCGACA TGGCAAGGAG GAGCTCGTTC CAGTCGTGTC 80 AGATAATATC CCTGTTCACT TTTGCCGTTG GAGTCAGTAT 120 CTGCTTAGGA TTCACTGCAC ATCGAATTAA GAGACACAC CTCCTGTTA TCAGTGCTT 240 GGATGGAGGA AAGGTCCTCC TACAGTGGTA TCAGACTCC 200 CCTGGACCAA CATCTCCGGA TCTTGCAGG GCAGGTGCTT 240 TGAACTTCAA GAGGCTGGAC CTCCTGATTG TCGCTGTGAC 280 AACTTGTGTA AGAGCTGAC CTCCTGATTG TCGCTGTGAC 280 TAAGAGCTGTC TTTGAAGACA GCCCCTGGCT CATGACTTTG 320 ATGAGCTGTC TTTGAAGACA GCCCCTGGCT GAGAATTAC CAGTTGCTGC CATGACTTTG 320 ATGAGCTGTC TTTGAAGACA GCCCCTGGCT GGGAGTGTAC 360 TGTCACTGCT CAGAGGACG CTCGCAGAATGA AGAAAATGCC 400 TGTCACTGCT CAGAGGACG CTTGGCCAGG GGAGACTGCT 440 GGTTGATGAT CCAAGTGGTT TCCAAAAGGAC AGTCGCATTG 480 GGTTGATGAT CCAAGTGGTT TCCAAAAGGAC ACTCTCCCTGATGC 520 CCTGCAGGGT TTTTTCCCC TCCATTAATC ATCTTCTCCG 560 TCGCATGGCTT CCCTGGATCA TACATGAAGA AAGACATGC 520 CCTGCAGGGT TTTTTCCCC TCCATTAATC ATCTTCTCCG 560		(2)	IN	FORM	OITA	N FO	r se	Q ID	NO:	68:				
(iii) HYPOTHETICAL: No  (ix) FEATURE: (A) NAME/KEY: A2058 ATX cDNA (B) LOCATION: (C) IDENTIFICATION METHOD: (D) OTHER INFORMATION:  (xi) SEQUENCE DESCRIPTION: SEQ ID NO:68:  25 CGTGAAGGCA AAGAGAACAC GCTGCAAAAG GCTTCCAAGA 40 ATCCTCGACA TGGCAAGAG GAGCTCGTTC CAGTCGTGTC 80 AGATAATATC CCTGTTCACT TTTGCCGTTG GAGTCAGTAT 120 CTGCTTAGGA TCACTGCAC ATCGAATAA GAGAGCAGAA 160 GGATGGGAGG AAGGTCCTC TACAGTGCTA TCAGACTCCC 200 CCTGGACCAA CATCTCCGGA TCTTGCAAGG GCAGGTGCTT 240 TGAACTTCAA GAGGCTGGAC CTCCTGATTG TCGCTGTGAC 280 ACTTGTGTA AGAGCTATAC CAGTTGCTGC CATGACTTTG 320 ATCGACTGTG TTTGAAGACA GCCCTGGCT GGGAGTTAC 360 TAAGGACAGA TGTGGAGAAG TCAGAAATGA AGAAAATGCC 400 TGTCACTGCT CAGAGGACTG CTTGGCCAGG GGAGACTGCT 440 GTACCAATTA CCAAGTGGTT TGCAAAGGAG AGTCGCATTG 480 GGTTGATGAT GACTGTGAGG AAATAAAAGGC CGCAGAATGC 520 CCTGCAGGGT TTGTTCGCCC TCCATTAATC ATCTTCTCCG 560 TGGATGGCTT CCGTGGCATCA TACATGAAGA AAGGCAGCAA 6000	15		(i	)	(A (B (C	) L ) T	ENGT YPE: TRAN	H: nu DEDN	3251 clei ESS:	c ac do	id uble			
(ix) FEATURE: (A) NAME/KEY: A2058 ATX cDNA (B) LOCATION: (C) IDENTIFICATION METHOD: (D) OTHER INFORMATION:  (Xi) SEQUENCE DESCRIPTION: SEQ ID NO:68:  25 CGTGAAGGCA AAGAGAACAC GCTGCAAAAG GCTTCCAAGA 40 ATCCTCGACA TGGCAAGGAG GAGCTCGTTC CAGTCGTGTC 80 AGATAATATC CCTGTTCACT TTTGCCGTTG GAGTCAGTAT 120 CTGCTTAGGA TTCACTGCAC ATCGAATTAA GAGAGCAGAA 160 GGATGGGAGG AAGGTCCTCC TACAGTGCTA TCAGACTCCC 200 CCTGGACCAA CATCTCCGGA TCTTGCAGTG TCAGACTCCC 200 CCTGGACCAA CATCTCCGGA TCTTGCAGG GCAGGTGCTT 240 AACTTGTGAA AGAGCATAAC CAGTGCTA TCAGACTCCC 280 AACTTGTGAACACA GAGCACAC CTCCTGATTG TCGCTGTGAC 280 AACTTGTGAACACA GCCCGTGGCT GGAAGTGTAC 360 TAAGGACAGA TGTGGGAGAG TCAGAAAATGA AGAAAATGCC 400 TGTCACTGCT CAGAGGACTG CTTGGCCAGG GGAGACTGCT 440 GGTTGATGAT CCAAGTGGTT TGCAAAGGAC AGTCGCATTG 480 GGTTGATGAT TTGTCGCCC TCCATTAATC ATCTTCTCCG 560 TCGCATGGCTT CCGTGCATCA TACATGAAGA AAGGCAGCAA 600			(i	i)	MC	LECU	LE I	YPE:	cD	NA				
(A) NAME/KEY: A2058 ATX CDNA  (B) LOCATION:  (C) IDENTIFICATION METHOD:  (D) OTHER INFORMATION:  (Xi) SEQUENCE DESCRIPTION: SEQ ID NO:68:  (XI) SEQUENCE DES			(i	ii)	HY	POTH	ETIC	AL:	No					
CGTGAAGGCA AAGAGAACAC GCTGCAAAAG GCTTCCAAGA ATCCTCGACA TGGCAAGAG GAGCTCGTTC CAGTCGTGTC AGATAATATC CCTGTTCACT TTTGCCGTTG GAGTCAGTAT CTGCTTAGGA TTCACTGCAC ATCGAATTAA GAGAGCAGAA GGATGGGAGG AAGGTCCTCC TACAGTGCTA TCAGACTCCC CCTGGACCAA CATCTCCGGA TCTTGCAAGG GCAGGTGCTT TGAACTTCAA GAGGCTGGAC CTCCTGATTG TCGCTGTGAC AACATTGTGTA AGAGCTATAC CAGTTGCTGC CATGACTTTG ATGAGCTGTG TTTGAAGACA GCCCGTGGCT GGGAGTGTAC ATGAGCAGAA TGTGGAGAAG TCAGAAATGA AGAAAATGCC TGTCACTGCT CAGAGGACTG CTTGGCCAGG GGAGACTGCT GTACCAATTA CCAAGTGGTT TGCAAAGGAG AGTCGCATTG GGTTGATGAT GACTGTGAGG AAATAAAGGC CGCAGAATGC CCTGCAGGGT TTGTTCGCCC TCCATTAATC ATCTTCTCCG TGGATGGCTT CCGTGCATCA TACATGAAGA AAGGCAGCAA 600	20		(i	.x)	(E (E	(a) N (b) I (c) I	IAME/ OCAT DENT	ION:	CATIC	N ME	THOL			
ATCCTCGACA TGGCAAGGAG GAGCTCGTTC CAGTCGTGTC AGATAATATC CCTGTTCACT TTTGCCGTTG GAGTCAGTAT CTGCTTAGGA TTCACTGCAC ATCGAATTAA GAGAGCAGAA 160 GGATGGGAGG AAGGTCCTCC TACAGTGCTA TCAGACTCCC 200 CCTGGACCAA CATCTCCGGA TCTTGCAAGG GCAGGTGCTT 240 TGAACTTCAA GAGGCTGGAC CTCCTGATTG TCGCTGTGAC 280 AACTTGTGTA AGAGCTATAC CAGTTGCTGC CATGACTTTG 320 ATGAGCTGTG TTTGAAGACA GCCCGTGGCT GGGAGTGTAC 360 TAAGGACAGA TGTGGAGAAG TCAGAAATGA AGAAAATGCC 400 TGTCACTGCT CAGAGGACTG CTTGGCCAGG GGAGACTGCT 440 GGTTGATGAT CCAAGTGGTT TGCAAAGGAG AGTCGCATTG 480 GGTTGATGAT GACTGTGAGG AAATAAAGGC CGCAGAATGC 520 CCTGCAGGGT TTGTTCGCCC TCCATTAATC ATCTTCTCCG 560 TGGATGGCTT CCGTGCATCA TACATGAAGA AAGGCAGCAA 600			()	ci)	SI	EQUE	ICE I	DESCF	RIPTI	ON:	SEQ	ID 1	10:68:	
ATCCTCGACA TGGCAAGGAG GAGCTCGTTC CAGTCGTGTC AGATAATATC CCTGTTCACT TTTGCCGTTG GAGTCAGTAT CTGCTTAGGA TTCACTGCAC ATCGAATTAA GAGAGCAGAA 160 GGATGGGAGG AAGGTCCTCC TACAGTGCTA TCAGACTCCC 200 CCTGGACCAA CATCTCCGGA TCTTGCAAGG GCAGGTGCTT 240 TGAACTTCAA GAGGCTGGAC CTCCTGATTG TCGCTGTGAC 280 AACTTGTGTA AGAGCTATAC CAGTTGCTGC CATGACTTTG 320 ATGAGCTGTG TTTGAAGACA GCCCGTGGCT GGGAGTGTAC 360 TAAGGACAGA TGTGGAGAAG TCAGAAATGA AGAAAATGCC 400 TGTCACTGCT CAGAGGACTG CTTGGCCAGG GGAGACTGCT 440 GGTTGATGAT CCAAGTGGTT TGCAAAGGAG AGTCGCATTG 480 GGTTGATGAT GACTGTGAGG AAATAAAGGC CGCAGAATGC 520 CCTGCAGGGT TTGTTCGCCC TCCATTAATC ATCTTCTCCG 560 TGGATGGCTT CCGTGCATCA TACATGAAGA AAGGCAGCAA 600	25	رروم	רכ <i>י</i> א א ר	מפרא	AACI	ממטע	CAC (	3 <b>ር</b> ፐርረ	ZAAAZ	AG GO	CTTCC	CAAGI	4	40
AACTTGTGTA AGAGCTATAC CAGTTGCTGC CATGACTTTG  ATGAGCTGTG TTTGAAGACA GCCCGTGGCT GGGAGTGTAC  TAAGGACAGA TGTGGAGAAG TCAGAAATGA AGAAAATGCC  TGTCACTGCT CAGAGGACTG CTTGGCCAGG GGAGACTGCT  GTACCAATTA CCAAGTGGTT TGCAAAGGAG AGTCGCATTG  GGTTGATGAT GACTGTGAGG AAATAAAGGC CGCAGAATGC  CCTGCAGGGT TTGTTCGCCC TCCATTAATC ATCTTCTCCG  TGGATGGCTT CCGTGCATCA TACATGAAGA AAGGCAGCAA  600		ATO AGA CTO GGA CCT	CCTC( ATAAT GCTT! ATGG( IGGA(	FACA FATC AGGA FAGG CCAA	TGG( CCT( TTC) AAG( CAT(	CAAG( STTC! ACTG( STCC' CTCC(	EAG (ACT TACE IN COLUMN TO TAC	FAGCT TTTG( ATCG) TACA( TCTT(	rcgt: Ccgt: Aatt: Gtgc: Gcaa(	rc ci rg gi AA gi ra to eg go	AGTC( AGTC) AGAG( CAGA( CAGG'	STGT( AGTA CAGA! CTCC( CGCT)		120 160 200 240
ATGAGCTGTG TTTGAAGACA GCCCGTGGCT GGGAGTGTAC 360 TAAGGACAGA TGTGGAGAAG TCAGAAATGA AGAAAATGCC 400 TGTCACTGCT CAGAGGACTG CTTGGCCAGG GGAGACTGCT 440 GTACCAATTA CCAAGTGGTT TGCAAAGGAG AGTCGCATTG 480 GGTTGATGAT GACTGTGAGG AAATAAAGGC CGCAGAATGC 520 CCTGCAGGGT TTGTTCGCCC TCCATTAATC ATCTTCTCCG 560 TGGATGGCTT CCGTGCATCA TACATGAAGA AAGGCAGCAA 600	30	TG/ AA(	AACT. CTTG:	rcaa rgta	AGA	GCTA'	FAC (	CAGT	rgct(	GC C	ATGA	CTTT(	3	320
TGGATGGCTT CCGTGCATCA TACATGAAGA AAGGCAGCAA 600	<i>3</i> 0	ATC TA TG GT. GG	GAGC' AGGA( TCAC' ACCA TTGA' TGCA	IGTG CAGA IGCT ATTA IGAT GGGT	TTTC TGTC CAG CCA GAC TTG	GAAGA GGAGA AGTG TGTG TTCG	ACA (AAG STT STEEL STT STEEL STT STT STT STT STT STT STT STT STT ST	GCCC TCAG CTTG TGCA AAAT TCCA	GTGG( AAAT( GCCA( AAGG AAAG TTAA	CT GO GA AO GG GO AG AO GC C TC A	GGAG' GAAA GAGA GTCG GCAG TCTT	IGTA ATGC CTGC CATT AATG CTCC	C C G G	400 440 480 520 560
	35	TG	CATG	GCTT	CCG	TGCA	TCA	TACA	TGAA	GA A	AGGC	AGCA	A	600 640

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	ር እ ርጥርጥር ርርጥ	<b>ACATGAGGCC</b>	GGTGTACCCA	ACTAAAACCT	680
	CHCICICCCI	ACAI GAGGCC	GCCACTGGGC	TATATCCAGA	720
	TICCIAACII	AIACACITIG	ATTCA ATCTA	TGATCCTGTA	760
	ATCACATOGA	CTTTTCATCT	GCGAGGGCGA	CACAAATTTA	800
	AMONDACATO	GTGGGGAGGT	CAACCCCTAT	GGATTACAGC	840
	ATCATAGATG	GGGGTGAAAG	CTCCD & CATT	CTTTTCTCC	880
	CACCAAGCAA	CTCACGAGCG	CIGGAACAII	ACCATATTCC	920
5	GTTGTCATCC	CICACGAGCG	CATCACACCC	CTTCGGTCTA	960
	GGTGGCTCAC	CCIGCCAGAI	CHIGAGAGGC	TGGACACAAA	1000
	TGCCTTCTAT	TCGGCCCTGA	CIGATITUTE	TOURCECTEAC	1040
	TATGGCCCTT	GGCTAAGAGA	GGAGAGIAGI	AACTTCCCCC	1080
	CTTTTACTCC	GGCIAAGAGA	CCIAAGAGGA	TCCAAAGAAA	1120
	TAAGAGGAGA	CAGGAAAGAC	CAGIIGCICC	TATGCTGCGG	1160
	AGAAGAAGAA	AAATACATAG	GATGGATCAT	TAIGCIGCGG	1200
10	AAACTCGTCA	GGACAAAATG	ACAAATCCIC	1 GAGGGAAA1	1240
10	CGACAAAATT	GTGGGGCAAT	TAATGGATGG	ACTGAAACAA	1240
	CTAAAACTGC	GTCGGTGTGT	CAACGTCATC	TTTGTCGGAG	1320
	ACCATGGAAT	GGAAGATGTC	ACATGTGATA	GAACTGAGII	1360
	CTTGAGTAAT	TACCTAACTA	ATGTGGATGA	TATTACTTTA	
	GTGCCTGGAA	CTCTAGGAAG	AATTCGATCC	AAATTTAGCA	1400
	ACAATGCTAA	ATATGACCCC	AAAGCCATTA	TTGCCAATCT	1440
	CACGTGTAAA	AAACCAGATC	AGCACTTTAA	GCCTTACTTG	1480
15				GCCAACAACA	1520
13	GAAGAATTGA	GGATATCCAT	TTATTGGTGG	AACGCAGATG	1560
	GCATGTTGCA	AGGAAACCTT	TGGATGTTTA	TAAGAAACCA	1600
	TCAGGAAAAT	GCTTTTTCCA	GGGAGACCAC	GGATTTGATA	1640
	ACAAGGTCAA	CAGCATGCAG	ACTGTTTTTG	TAGGTTATGG	1680
	CCCAACATTT	AAGTACAAGA	CTAAAGTGCC	TCCATTTGAA	1720
	AACATTGAAC	TTTACAATGT	TATGTGTGAT	CTCCTGGGAT	1760
	TGAAGCCAGC	TCCTAATAAT	GGGACCCATG	GAAGTTTGAA	1800
20	TCATCTCCTG	CGCACTAATA	CCTTCAGGCC	AACCATGCCA	1840
	GAGGAAGTTA	CCAGACCCAA	TTATCCAGGG	ATTATGTACC	1880
	TTCAGTCTGA	TTTTGACCTG	GGCTGCACTT	GTGATGATAA	1920
	GGTAGAGCCA	AAGAACAAGT	TGGATGAACT	CAACAAACGG	1960
	CTTCATACAA	AAGGGTCTAC	AGAAGAGAGA	CACCTCCTCT	2000
	ATGGGCGACC	TGCAGTGCTT	TATCGGACTA	GATATGATAT	2040
	CTTATATCAC	ACTGACTTTG	AAAGTGGTTA	TAGTGAAATA	2080
25				GTTTCCAAAC	2120
2.5	AGGCTGAGGT	TTCCAGCGTT	CCTGACCATC	TGACCAGTTG	2160
	CGTCCGGCCT	GATGTCCGTG	TTTCTCCGAG	TTTCAGTCAG	2200
	AACTGTTTGG	CCTACAAAAA	TGATAAGCAG	ATGTCCTACG	2240
	GATTCCTCTT	TCCTCCTTAT	CTGAGCTCTT	CACCAGAGGC	2280
	TAAATATGAT	GCATTCCTTG	TAACCAATAT	GGTTCCAATG	2320
	TATCCTGCTT	TCAAACGGGT	CTGGAATTAT	TTCCAAAGGG	2360
	TATTGGTGAA	GAAATATGCT	TCGGAAAGAA	ATGGAGTTAA	2400
30	CGTGATAAGT	GGACCAATCT	TCGACTATGA	CTATGATGGC	2440
	TTACATGACA	CAGAAGACAA	AATAAAACAG	TACGTGGAAG	2480
	GCAGTTCCAT	TCCTGTTCCA	ACTCACTACI	ACAGCATCAT	2520
	CACCAGCTGT	CTGGATTTCA	CTCAGCCTGC	CGACAAGTGT	2560
				CTGCCTCACC	2600
	GGCCTGACAA	CGAGGAGAGC	TGCAATAGC	CAGAGGACGA	2640
				GCACACAGCT	2680
25				CTGGACTTCT	2720
35	TCCGAAAGAC	CAGCCGCAGC	TACCCAGAA	1 TCCTGACACT	2760

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5	CAAGACATAC CTGCATACAT ATGAGAGCGA GATTTAACTT TCTGAGCATC TGCAGTACAG TCTTATCAAC TGGTTGTATA TTTTTATATT GTTTTTGTAT TTATTAATTT GAAACCAGGA CATATAAAAAT GTTAGTATTT TAATCCTGTA CCAAATCTGA CATATTATGC CTGAATGACT CCACTGTTTT TCTCTAATGC TTGATTTAGG TAGCCTTGTG TTCTGAGTAG AGCTTGTAAT AAATACTGCA GCTTGAGAAA AAGTGGAAGC TTCTAAATGG TGCTGCAGAT TTGATATTTG CATTGAGGAA ATATTAATTT TCCAATGCAC AGTTGCCACA TTTAGTCCTG TACTGTATGG AAACACTGAT TTTGTAAAGT TGCCTTTATT TGCTGTTAAC TGTTAACTAT GACAGATATA TTTAAGCCTT TAAAAAAAAAA	2800 2840 2880 2920 2960 3000 3040 3120 3160 3200 3240 3251
10		
15	(2) INFORMATION FOR SEQ ID NO:69:  (i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 915  (B) TYPE: amino acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: Unknown	·
	(ii) MOLECULE TYPE: cDNA	
	·	
	(iii) HYPOTHETICAL: No	
20	<pre>(ix) FEATURE:     (A) NAME/KEY: A2058 ATX protein     (B) LOCATION:     (C) IDENTIFICATION METHOD:     (D) OTHER INFORMATION:</pre>	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:69:	
25	Met Ala Arg Arg Ser Ser Phe Gln Ser Cys Gln Ile	
- <del>-</del>	1 5 10 Ile Ser Leu Phe Thr Phe Ala Val Gly Val Ser Ile 15 20	
	Cys Leu Gly Phe Thr Ala His Arg Ile Lys Arg Ala 25 30 35	
	Glu Gly Trp Glu Glu Gly Pro Pro Thr Val Leu Ser 40 45	
30	Asp Ser Pro Trp Thr Asn Ile Ser Gly Ser Cys Lys 50 55 60	
	Gly Arg Cys Phe Glu Leu Gln Glu Ala Gly Pro Pro 65 70	
	Asp Cys Arg Cys Asp Asn Leu Cys Lys Ser Tyr Thr 75 80	
35	Ser Cys Cys His Asp Phe Asp Glu Leu Cys Leu Lys 85 90 95	

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•	Thr	Ala	Arg	Gly 100	Trp	Glu	Cys	Thr	Lys 105	Asp	Arg	Cys
	Gly	Glu 110	Val	Arg	Asn	Glu	Glu 115	Asn		Cys	His	Cys 120
	Ser		Asp	Cys	Leu 125	Ala		Gly	Asp	Cys 130	Cys	Thr
5	Asn	Tyr	Gln 135	Val		Cys	Lys	Gly 140	Glu	Ser	His	Trp
3	Val 145	Asp		Asp	Cys	Glu 150	Glu	Ile	Lys	Ala	Ala 155	Glu
	Cys	Pro	Ala	Gly 160	Phe	Val	Arg	Pro	Pro 165	Leu	Ile	Ile
		170					175				Met	180
10	-	_		_	185					190	Lys	
			195	_				200			Arg	
	205	_				210					Tyr 215	
				220					225		Gly	
15		230					235				Asp	240
					245					250	Asn	
			255					260			Thr	
20	265	_		_		270					Phe 275	
20				280					285		Leu	
		290					295				Glu	300
					305					310		
25			315					320			Pro	Lys
	325					330					335	Gln
	_			340	_				345			Arg
		350					355					360 Thr
30	_				365					370		
	_		375	_				380	)			Ile
	385	;				390	)				395	
				400	)				405	5		Phe
35	val	410	_	nls	, сту	Met	415	_	, Agl	. 1111	. cys	Asp 420

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					Leu 425								
			425		Leu			44.0					
	–		Ser		Phe	4511							
5	Pro			160	Ile				400				
		470		His	Phe		4/2						
		Pro	Lys		Leu 485					せりい	,		
			405		His	Leu		ついい					
10			Ala	Arg	Lys	<b>~ 1 1</b>							
	Pro	Sei		E 27 /	Cys				: 3 Z	,			
		E 2 (	`		val		5.35	)					
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20			^		e Arg		~ ~	~					
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Leu Phe Pro Pro Tyr Leu Ser Ser Ser Pro Glu Ala
             735
                                 740
     Lys Tyr Asp Ala Phe Leu Val Thr Asn Met Val Pro
                         750
     Met Tyr Pro Ala Phe Lys Arg Val Trp Asn Tyr Phe
                 760
                                      765
     Gln Arg Val Leu Val Lys Lys Tyr Ala Ser Glu Arg
                             775
5
     Asn Gly Val Asn Val Ile Ser Gly Pro Ile Phe Asp
                     785
     Tyr Asp Tyr Asp Gly Leu His Asp Thr Glu Asp Lys
             795
                                 800
     Ile Lys Gln Tyr Val Glu Gly Ser Ser Ile Pro Val
                         810
     Pro Thr His Tyr Tyr Ser Ile Ile Thr Ser Cys Leu
10
                 820
                                      825
     Asp Phe Thr Gln Pro Ala Asp Lys Cys Asp Gly Pro
                             835
     Leu Ser Val Ser Ser Phe Ile Leu Pro His Arg Pro
                     845
     Asp Asn Glu Glu Ser Cys Asn Ser Ser Glu Asp Glu
                                  860
     Ser Lys Trp Val Glu Glu Leu Met Lys Met His Thr
15
                         870
     Ala Arg Val Arg Asp Ile Glu His Leu Thr Ser Leu
                 880
                                      885
     Asp Phe Phe Arg Lys Thr Ser Arg Ser Tyr Pro Glu
                              895
     Ile Leu Thr Leu Lys Thr Tyr Leu His Thr Tyr Glu
                     905
20
     Ser Glu Ile
             915
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DMCDCCID: -\MC 953222142 L

#### CLAIMS:

- 1. A DNA segment coding for a polypeptide comprising an amino acid sequence corresponding to autotaxin, or a fragment thereof having at least 5 amino acids.
- 5 2. The DNA segment according to claim 1, wherein said DNA segment encodes the amino acid sequence selected from the group consisting of SEQ ID NO:1 through SEQ ID NO:11, SEQ ID NO:26 through SEQ ID NO:34, SEQ ID NO:36, SEQ ID NO:38 through SEQ ID NO:52, SEQ ID NO:66
  10 and SEQ ID NO:69.
  - 3. An isolated polypeptide comprising an amino acid sequence corresponding to autotaxin, or a fragment thereof having at least 5 amino acids.
- 4. The polypeptide according to claim 3, wherein said amino acid sequence comprises the amino acid sequence selected from the group consisting of SEQ ID NO:1 through SEQ ID NO:11, SEQ ID NO:26 through SEQ ID NO:34, SEQ ID NO:36, and SEQ ID NO:38 through SEQ ID NO:52, SEQ ID NO:67 and SEQ ID NO:69.
- 5. An isolated polypeptide bound to a solid support, comprising an amino acid sequence corresponding to autotaxin, or a fragment thereof having at least 5 amino acids.
- 6. The polypeptide according to claim 5,
  wherein said polypeptide comprises the amino acid sequence
  selected from the group consisting of the SEQ ID NO:1
  through SEQ ID NO:11, SEQ ID NO:26 through SEQ ID NO:34,
  SEQ ID NO: 36, SEQ ID NO:38 through SEQ ID NO:52., SEQ ID
  NO:67 and SEQ ID NO:69.
- 7. A recombinant DNA molecule comprising a vector and the DNA segment according to claim 1.
  - 8. A cell that contains the recombinant DNA molecule according to claim 7.
- An antibody having binding affinity for
   autotaxin, or binding fragment thereof.

10. A method of producing a recombinant autotaxin polypeptide said method comprising:

culturing a cell containing the recombinant DNA molecule of claim 7 under conditions such that the DNA

segment is expressed, producing said polypeptide; and isolating said polypeptide.

11. A method of purifying the autotaxin peptide of claim 3, comprising the steps of:

 i) collecting and concentrating supernatant from cultured A2058 human melanoma cells whereby a first preparation of said peptide is produced;

ii) salt fractionating said first preparation to produce a second peptide preparation;
iii) isolating said peptide from said second preparation so that said peptide is obtained in substantially pure form.

12. The method of claim 11, wherein said isolating step is effected by column chromatography.

13. An isolated DNA encoding an autotaxin protein or fragment thereof wherein said DNA includes a nucleic acid sequence selected from the group consisting of SEQ ID NO:35, SEQ ID NO:37 and SEQ ID NO:38.

14. The DNA segment according to claim 1, wherein said DNA fragment comprises any one of the SEQ ID NO:12 through SEQ ID NO:25, or SEQ ID NO:39 through SEQ ID NO:52.

15. The DNA segment according to claim 13 wherein said DNA segment comprises any one of the SEQ ID NO:12 through SEQ ID NO:25.

16. An isolated polypeptide comprising an amino acid sequence corresponding to autotaxin.

17. A polypeptide bound to a solid support and comprising an amino acid sequence corresponding to autotaxin.

18. A recombinant autotaxin polypeptide

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according to claim 3.

19. An isolated polypeptide according to claim
3 having cell motility activity.

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FIG. 1

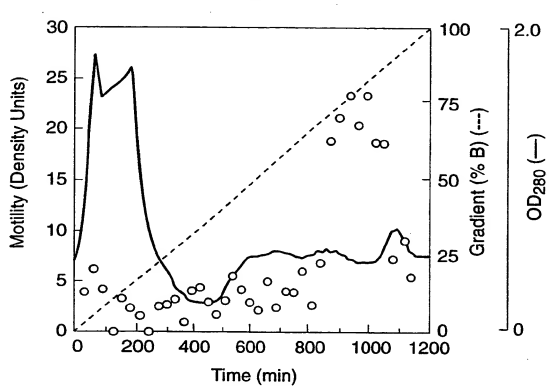
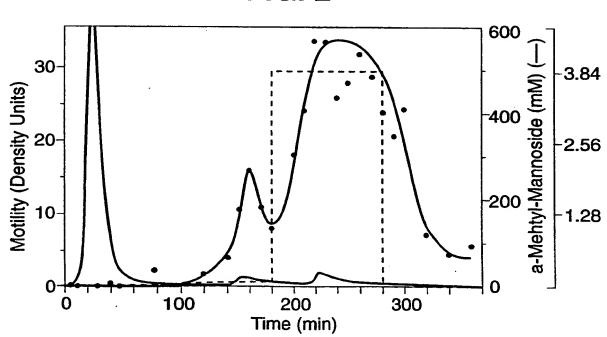


FIG. 2



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FIG. 3

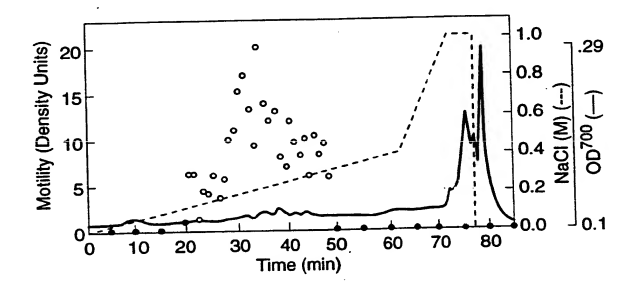


FIG. 4 0.32 30 Motility (Density Units) 15 0.15 10 5 0 100 110 90 80 50 70 60 Time (min)

FIG. 5 25 1.0 0.04 Motility (Density Units) 20 0.8 0.03 NaCl (M) (--) 15 0.02 10 0.01 5 0.2 0.00 0 0.0 10 ە<del>رە</del> 60 50 Ó 20 30 40 70 Time (min)

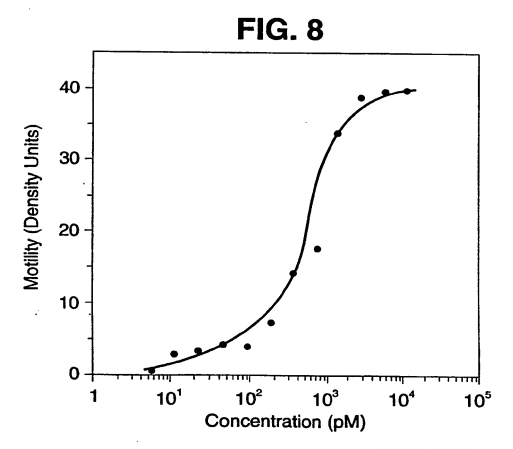
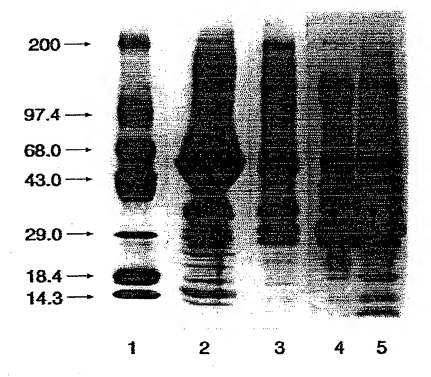
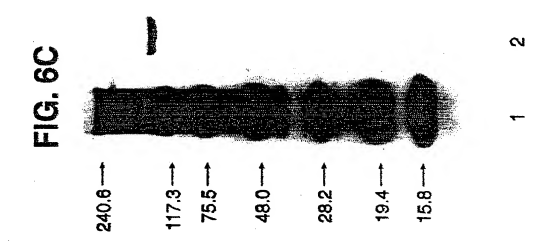


FIG. 6A





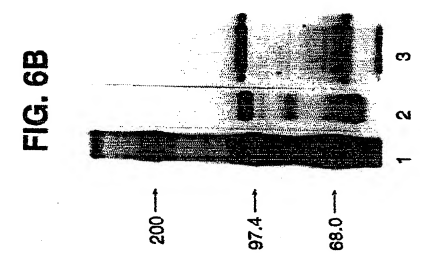


FIG. 7

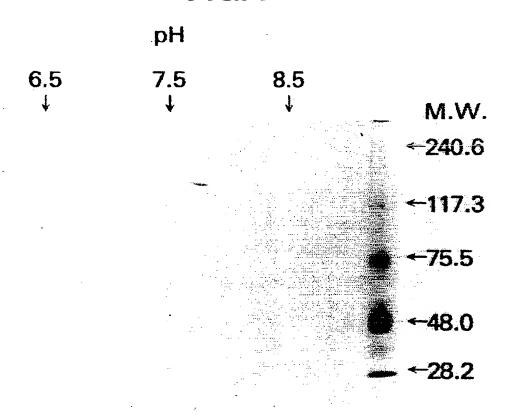


FIG. 16

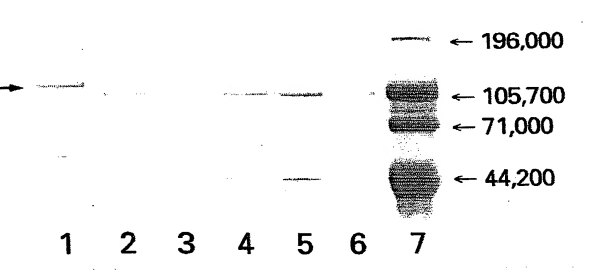




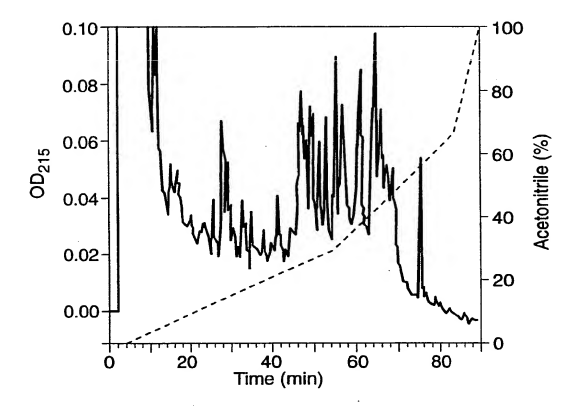
FIG. 10

Lower Walls

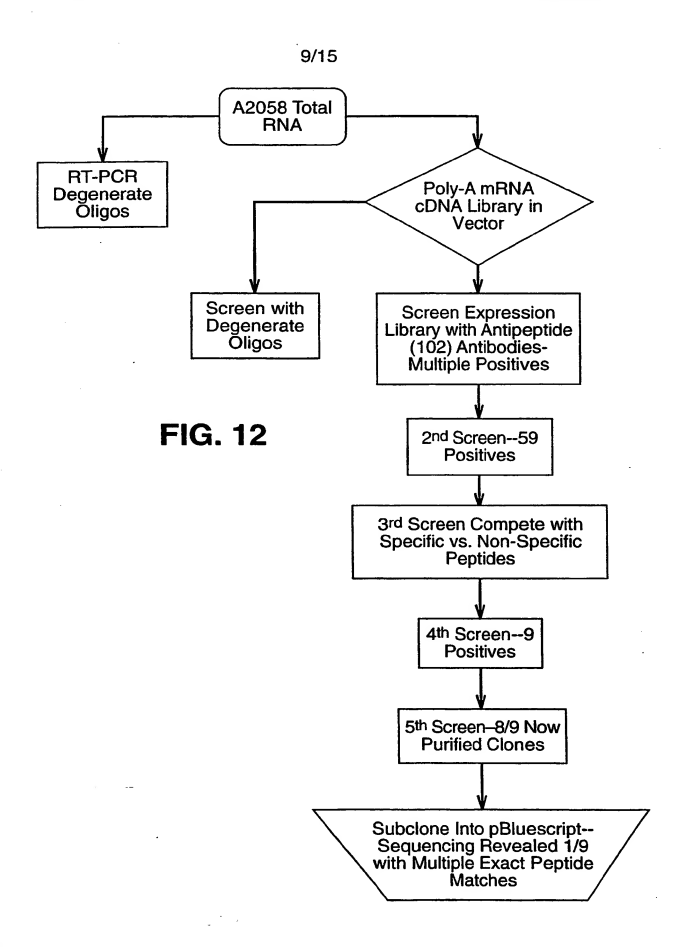
#### **Upper Walls**

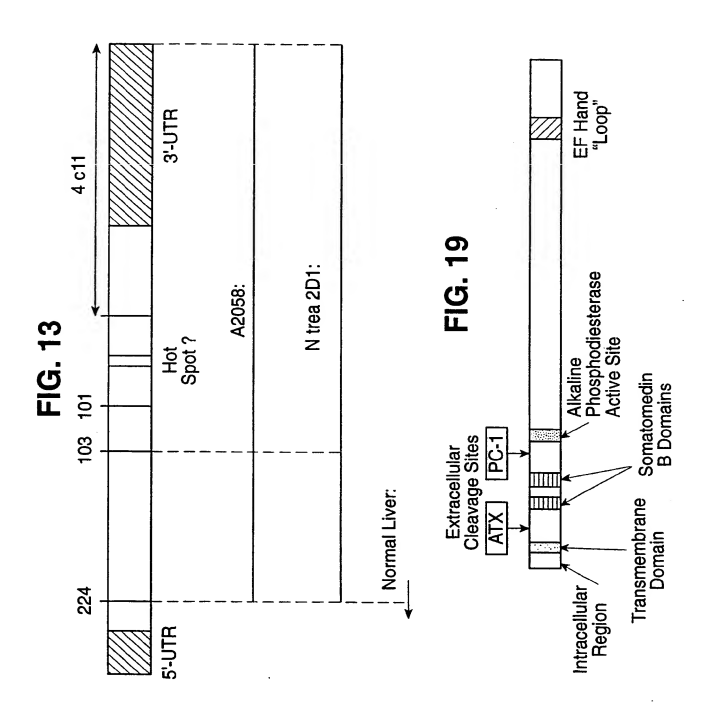
	0	0.01%	0.1%
0	4.8 ± 0.3	13.7 ± 0.8	33.8 ± 1.6
0.01%	45.4 ± 4.0	39.3 ± 2.6	34.9 ± 1.4
0.1%	75.6 ± 1.8	58.3 ± 3.1	41.0 ± 3.4

FIG. 11

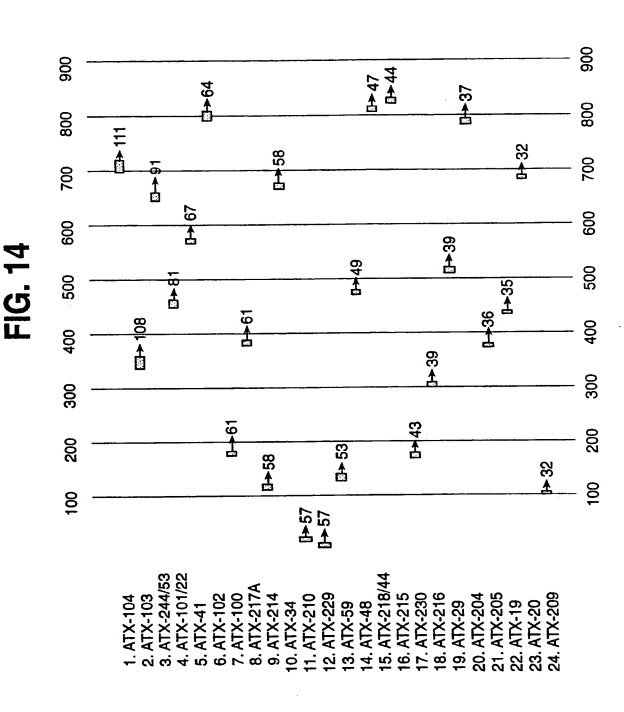


WO 95/32221 PCT/US95/06613

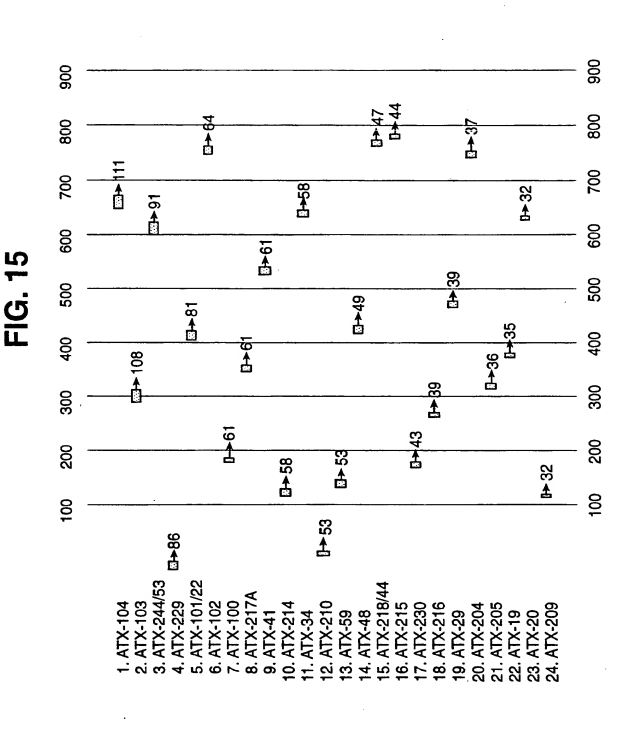




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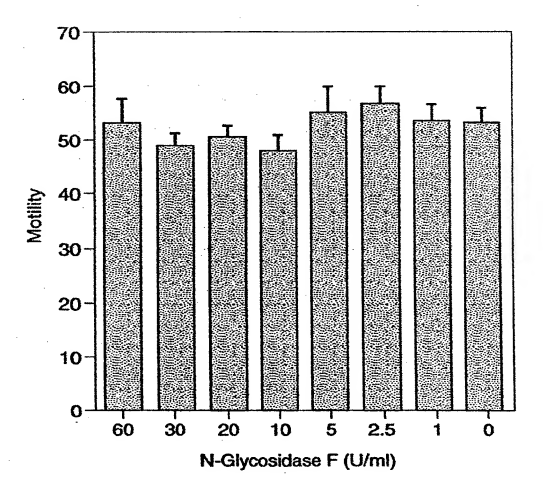
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**FIG. 17A** 

<--125 kDa

### FIG. 17B



SUBSTITUTE SHEET (RULE 26)

# FIG. 18A

hATX	MARRSSFQSCQIISLFTFAVGVSICLGFTAHRIKRAEGWEEGPPTVLSDSPWTNISG
hPC1	MDVGEEPLEKAARARTAKDPNTYKVLSLVLSVCVLTTILGCIFGLKPSCAKEVK.SCKGRCFERTFGNCRCDAACVELGHCCLDY 84
hATX	DELCLKTARGWECTKDRCGEVRNEENA
hPCl	
hATX	
hPC1	
hATX	VVIPHERRILTILRWLTLPDHERPSVYAFYSEQPDFSGHKYGPFGPEESSYGSPFTPAKRPKRKVAPKRRQERPVAPPKKRRKIHRMDHYAAET 372
hPC1	
hATX	RQDKMTNPLREIDKIVGQLMDGLK
hPC1	
hATX	QHFKPYLKQHLPKRLHYANNRRIEDIHLLVERRWHVARKPLDVYKKPSGKCFFQGDHGFDNKVNSMQTVPVGYGPTFKYKTKVPPFENIELYNVMCDLIG 570
hPC1	QHFKPYLKHGLPKRLHFAKSDRIEPLTFYLDPQWQLALNPSERKYCGSGFHGSDNVFSNMQALFVGYGPGFKHGIEADTFENIEVYNLMCDLLN 526

## FIG. 18B

4 4 4 4 4 4 4 4	hatx LKP hpcl LTP hatx DFE hatx FQF hpcl FHI hpcl FHI hatx SKV	
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